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  JOURNAL OF ORGANIC CHEMISTRY, vol. 41,
  no. 12, 11th June 1976, pp. 2098-2102,
  Washington D.C. (US); D. JOHN ABERHART et
  al.: "Studies on the adduct of 4-phenyl-1,2,4triazoline-3,5-dione with Vitamin D3"

CHEMICAL ABSTRACTS, vol. 92, no. 19, 12th May 1980, p. 624, no. 164161d, Columbus, Ohio (US); E.ZBIRAI et al.: "Structural transformations on vitamin D"

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## D scription

This invention relates to novel intermediates in the productien of vitamin D analogues and of vitamin D analogues which may be produced therefrom.

In the past modified vitamin D derivatives have been prepared through modification of sterol precursors which are then converted intervitation D derivatives through a standard series of steps, normally preliminary conversions of  $\Delta^{5,7}$  dienes followed by irradiation of the dienes to give D vitamins. These procedures have serious flaws. First, all of the available methods for the synthesis of  $\Delta^{5,7}$  dienes tend to give mixtures of products or require a number of steps and proceed in relatively low yield. The second difficulty is that the only known transformation of the 5,7 dienes into the vitamins involves irradiation followed by thermal equilibration. Irradiation intrinsically gives rise to a mixture of byproducts. This limits the yield of the desired vitamin and furthermore complicates its recovery in pure form.

Previous attempts to modify the 17-side chain of vitamin D compounds have been unsuccessful due t instability problems. We have now found that vitamin  $D_2$  and related compounds can be converted to a protected form capable of withstanding the reaction conditions necessary for oxidative cleavage of the 22,23-double bond to form a 22-aldehyde which can then be converted to other derivatives as described hereinafter. In particular, we have found that vitamin  $D_2$  compounds in either the *cis* or *trans* configuration can be stabilised by formation of a Diels Alder dienophile adduct which can subsequently be reconverted to the *trans* form of the vitamin after the side-chain modification. The *trans* vitamin analogues can then be efficienctly converted into the active *cis* form by known reactions.

The formation of certain dienophile adducts of vitamin  $D_3$  has been described in the literature. D. J. Aberhart and A. Chi Tung Hsu (J. Org. Chem. Vol. 41, No. 12, 1976, 2098—2102) have described the formation of an adduct with 4-phenyl-1,2,4-triazoline-3,5-dione and E. Zbiral and W. Reischl (Proc. Workshop Vitamin D 1979, 4th. (Vitam. D. Basic Res. Its Clin Appl.), 21—24) have also described the adduct with sulphur dioxide. However, there is no disclosure of the use of these dienophiles to form adducts with vitamin  $D_2$  followed by oxidative cleavage at the 22,23-double bond while leaving double bonds in the 5,10-and 7,8-positions intact.

According to one feature of the present invention we provide compounds of the general formula I,

wherein R represents a hydrogen atom or a hydroxyl protecting group, Y represents a hydrogen atom or an optionally protected hydroxyl group, X represents — $SO_2$  or the residue of a diacylazo dienophile and either R¹ represents a halogen atom a hydrocarbylsulphonyloxy group or a group of the formula —Z—R³ (in which Z represents —O—, —S—, —SO—, —NR⁴— or —CR⁴R⁵— and R³, R⁴ and R⁵, which may be the same or different, each represents a hydrogen atom or a straight or branched aliphatic group having 1—12 carbon atoms and which may optionally carry one or more substituents) and R² represents a hydrogen atom or R¹ and R² together represent an oxo group or an optionally substituted alkylidene group, except that R¹ together with the group —CH(CH₃)CH to which they are attached do not represent a group having the branched 17β-hydrocarbyl side chain skeleton of vitamin D₂ or vitamin D₃.

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The abov compounds ar us ful intermediates in the preparation of vitamin D analogues i.e. comp unds of general formula IV and IVa

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wherein R, Y, R<sup>1</sup> and R<sup>2</sup> are as hereinbefore defined. The above compounds of general formulae IV and IVa are also novel and constitute a still further feature of this invention.

The use of the compounds of general formula I in the preparation of the novel compounds of formulae IV and IVa is illustrated in the reaction scheme of the accompanying drawings, R, Y, X, R<sup>1</sup> and R<sup>2</sup> being as defined above. The compounds of formula I—IV may also carry further groupings.

It should be noted that the Diels Alder adduct formed from either the 5,6-cis- or the 5,6-trans-vitamin starting material exists as a mixture of two possible isomers at the 6-position. However, since the eventual removal of the Diels Alder residue always generates a compound of the 5,6-trans configuration, there is no need to distinguish between such 6-isomers or to effect their separation.

We have found that using the above procedure a wide range of groups R1 may be introduced into the vitamin D structure. Thus, as indicated above R1 may be a group of the formula Z—R3, where Z is —0—, —S—, —SO—, —NR<sup>4</sup> or —CR<sup>4</sup>R<sup>5</sup>— and R<sup>3</sup>, R<sup>4</sup> and R<sup>5</sup>, which may be the same or different, are each a hydrogen atom or a straight or branched aliphatic group having 1-12 carbon atoms which may carry one or more substituents such as, for example halogen atoms (e.g. fluorine), or optionally protected hydroxyl groups.

In general it is preferred that the group R3 in the final products should be of the formula

(in order to provide a 17β-side chain of approximately the shape present in natural vitamin D compounds) with the possibility of substitution as described above. The heteroatoms Z, where present, do not greatly change the overall shape of the side chain.

In particular, the invention enables compounds of formula IV and IVa to be prepared in which R1 is of formula

wherein Z' represents -O--, -S--, -NH-- or -SO-- and R<sup>6</sup> represents a hydrogen atom or a hydroxyl protecting group, the 1a-position optionally carrying a hydroxyl or protected hydroxyl group, these being analogues of th active m tabolite 25-hydr xy vitamin D3.

Protected hydroxyl groups may, for example, be acyl groups e.g. alkanoyl groups (preferably having 1—6 carbon atoms), aralkanoyl groups (preferably having 7—15 carb in atoms), aroyl groups (preferably having 6-12 carbon atoms), cyclic ether groups or tri-hydrocarbylsilyl groups. Examples of such gr ups include acetyl, propionyl, benzoyl and tetrahydropyranyl groups and trihydrocarbylsilyl groups having up to three  $C_{1-6}$  alkyl,  $C_{6-12}$  aryl and/or  $C_{7-15}$  aralkyl groups.

The new synthetic analogues of the invention have modified vitamin D properties of int  ${\bf r}$  st in  ${\bf m}$  dicin .

The compounds of formulae IV and IVa in which R¹ has the abov meanings may be prepared, inter alia, by nucleophilic substitution reactions on compounds of f rmula IV and IVa in which R¹ repr sents a halog n at m, such as a chlorine, bromine or iodine atom, or a leaving group, for example a hydrocarbylsulphonyloxy group O—SO₂R² in which R² represents, for xample, an alkyl group (preferably having 1—6 carbon atoms), an aryl group (preferably having 7—15 carbon atoms). The tosyloxy group is preferred. Alternatively, the above compounds may be prepared from corresponding compounds of formula I and the dienophile group X removed subsequently. Since, however, the nucleophilic substitution reactions are mostly carried out in the presence of a base and since the protected compounds of formula I are less stable to base than the parent trienes of formula IV and IVa, the latter are commonly preferred substrates

In the formation of 22-thia compounds (in which Z is —S—), the nucleophilic reagent is conveniently the thiol of formula R³XX reacted in an inert solvent such as tetrahydrofuran in the presence of a non-nucleophilic base, for example an inorganic base such as sodium hydride or an organic base such as pyridine.

The corresponding sulphoxides (Z= —SO—) may be prepared by oxidation of the thia-compound (Z= —S—), for example using a peracid or salt as oxidising agent, e.g. a periodate. Mixtures of the (R) and (S) sulphoxides are normally formed and the invention extends to these separately and in admixture.

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The 22-oxa compounds of formula IV or IVa may be prepared by reaction of an alcohol of formula I, IV or IVa in which R¹ is OH, with an alkylating agent or alternatively by reaction of a reactive derivative thereof, with an alcoholate; these reactions are followed by deprotection when a compound of formula I is used. The reactive derivative may, for example, be a halide such as an iodide, or a hydrocarbylsulphonyloxy derivative, such as a tosyloxy derivative, the alcoholate being, for example, an alkali metal or thallium alcoholate of the alcohol R³OH. It is preferred, however, to react the compound of formula I, IV or IVa in which R¹ in which R¹ is OH with an epoxide. This generates a side chain carrying a hydroxyl group derived from the epoxide oxygen. Where it is desired to make 25-hydroxy-22-oxa vitamin D₃ derivatives, a suitable reagent is isobutylene epoxide.

The reaction is advantageously effected in an inert solvent, e.g. a hydrocarbon solvent such as benzene, in the presence of a non-nucleophilic base, conveniently an alkali metal t-alkoxide in the presence of a phase transfer agent such as a crown ether. Under such basic conditions, we have found it especially preferred to effect the reaction on a starting compound of formula IV or IVa, since the trienes are, as indicated above, more stable to these conditions than the dienophile-protected compounds of formula I.

The 22-aza compounds of formula I, IV or IVa may be prepared by reaction of a reactive derivative of an alcohol of formula I, IV or IVa in which R<sup>1</sup> is OH, for example a halide such as an iodide, or a hydrocarbylsulphonyloxy derivative such as a tosyloxy derivative, with an amine of formula R<sup>3</sup>R<sup>4</sup>NH. Due to the basicity of the reagent, a substrate of formula IV or IVa is preferred. Where the amine is liquid it is preferably reacted without added solvent.

The 22-aza derivatives may often conveniently be isolated as N-acylates, such as N-acetates, which may be formed by reaction with an appropriate acid anhydride.

The 22-hydrocarbylsulphonyloxy derivatives of formulae I, IV and IVa used in the above reactions may be prepared by reacting the corresponding alcohol with the appropriate hydrocarbylsulphonyl halide, e.g. tosyl chloride in the presence of a base such as pyridine. Best results have been obtained by effecting this reaction on a compound of formula I in which X is SO<sub>2</sub>, and removing the SO<sub>2</sub> residue by thermolysis, as described hereinafter.

The compounds of formula I, IV or IVa in which Z in R¹ is CR⁴R⁵ may be prepared by reacting compounds of formula I, IV or IVa carrying a hydrocarbylsulphonyloxy group R¹, e.g. a tosyl group, with carbon nucleophiles. Suitable carbon nucleophiles are Grignard reagents reacted in the presence of a copper catalyst, e.g. a cuprous salt. Thus, for example, 25-hydroxy vitamin D₃ and the 1α-hydroxy derivative thereof may be prepared by reacting an appropriate hydrocarbylsulphonyloxy derivative of formula I, IV or IVa with a Grignard reagent of the formula

(where R<sup>6</sup> has the above meaning) in tetrahydrofuran in the presence of cuprous iodide.

For the production of an alcohol of formula I in which R<sup>1</sup> is OH, for use in the preparation of the above novel vitamin D derivatives, the formyl group in the corresponding aldehyde of formula I (wherein R<sup>1</sup> and R<sup>2</sup> together represent oxo) must be reduced.

We have found that this can be effected readily, often in essentially quantitative yield, by reaction with a metal hydride reducing agent such as an alkali metal borohydride, e.g. sodium borohydride. It is noteworthy that this reduction retains the original configuration at the 20-carbon atom. Such alcohols ar also new compounds.

Compounds of formula I, IV or IVa may also be prepared in which R<sup>1</sup> is a divalent alkylidene group, which may carry substituents as described above for R<sup>3</sup>. Thus, for example, the aldehyde of formula I (wh rein R<sup>1</sup> and R<sup>2</sup> together repres nt oxo) may be reacted with an ylid, for example a Wittig r agent which may be represented by the general f rmula

$$(R^8)_3 P = R^{1A}$$

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wherein the groups  $R^8$ , which may be the same or different, are alkyl (preferably  $C_{1-8}$ ), aralkyl (preferably  $C_{7-15}$ ) or aryl (preferably  $C_{8-12}$ ) groups and  $R^{1A}$  is an alkylidene group (preferably having 1 to 8 carbon atoms) and may carry substituents as described for  $R^3$  above.

The Wittig reagent will normally be formed *in situ* by reaction of a quaternary salt thereof with a strong base in an inert solvent. Suitable bases include hydrocarbyl lithium compounds such as phenyl lithium and *n*-butyl lithium. Suitable solvents include ether solvents such as tetrahydropyran and diethyl ether. The aldehyde of formula I is preferably added immediately after the Wittig reagent has been formed.

The phosphonium salt precursor of the appropriate Wittig reagent for formation of the correct  $17\beta$ -side chain of 25-hydroxy vitamin  $D_3$  may, for example be prepared by reaction of isobutylene epoxide with methylenetriphenylphosphorane; the initially formed product in which  $R^5$  is H may if desired be protected, for example by formation of a tetrahydropyranyl or trihydrocarbylsilyl derivative. The phosphorane is preferably prepared by reaction of methyltriphenylphosphonium bromide in a cyclic ether solvent such as tetrahydrofuran in the presence of a strong base such as phenyl or n-butyl lithium, the isobutylene epoxide then being reacted in situ with a second equivalent of base. We have found the phosphonium bromide initially produced to be difficult to isolate and purify but that conversion to a tetraphenylborate salt enabled a relatively pure product to be obtained.

If it is desired to form a saturated side-chain, selective reduction of the newly formed 22,23-double bond is required. This was unexpectedly found to be possible using hydrogenation over 5% palladium on charcoal. It is noteworthy that this reduction restores the desired, "natural" configuration at the 22- and 23-carbon atoms. This route thus provides a further method of preparing compounds of formula I, IV or IVa in which R<sup>1</sup> is —CR<sup>3</sup>R<sup>4</sup>R<sup>5</sup>, as defined above.

It will be seen that the compounds of formula I in the above reaction scheme are key intermediates in the production of the new vitamin D analogues according to the invention. By way of illustration their preparation is now described in detail starting from vitamin D<sub>2</sub> or its 5,6-trans isomer.

The compound of formula III may be prepared by reaction of a vitamin D<sub>2</sub> compound of formula IIa or IIb with SO<sub>2</sub> or a diacylazo dienophile whereby the desired divalent grouping X is introduced.

Preferred diacylazo dienophiles are cyclic azo compounds such as phthalazine diones or triazoline diones; in general these may be represented by the formula V,

where W is a divalent aromatic carbocyclic group such as a 1,2-phenylene group or a group

where R<sup>9</sup> is an aryl group such as a phenyl group. The divalent aromatic group or the aryl group R<sup>9</sup> may carry substituents, for example C<sub>1-6</sub> alkyl or alkoxy groups, halogen atoms or nitro groups. Derivatives of formula III in which X is of formula V are also new compounds.

Where the dienophile is SO<sub>2</sub>, this may simply be reacted with the vitamin D<sub>2</sub> compound in a suitable solvent, for example aqueous media capable of dissolving the vitamin. A well stirred mixture of water and a hydrocarbon solvent such as benzene is particularly useful. Basic conditions are preferably used, e.g. using an inorganic base such as an alkali metal bicarbonate. Where the dienophile is a cyclic azo compound of formula V in which W is a group

this may be added to the starting vitamin D<sub>2</sub> compound in solution in a suitable solvent such as thyl acetat . Where W is a divalent-1,2-arylene grouping as in phthalazine 1,2-dione, however, this is preferably formed *in situ* by oxidation of the corresponding cyclic hydrazide, e.g. phthalhydrazide. Thus the vitamin D<sub>2</sub>

compound may be reacted in solution in an inert solvent such as a halogenated hydrocarbon with the cyclic hydrazide in the presence of an oxidising agent such as lead tetraacetate/ac tic acid.

After formation of the adduct of formula III, the 22,23-double bond may be cleaved to form the 22-formyl derivative of formula I by known oxidative techniques such as permanganate/periodate, osmate/periodate or, most pref rably oz n lysis. W have found that this reaction proceeds selectively in high yield with little cleavage of the 7,8-double b nd and, in particular, with no disturbance of the stereochemistry at the 20-position.

Oznolysis may be effected by passing ozone, preferably diluted with a further gas such as oxygen, through a solution of the compound of formula III in a solvent therefor to form an ozonide which is then reductively cleaved by a suitable reducing agent. A suitable solvent is, for example, a halogenated hydrocarbon such as dichloromethane, a ketone, e.g. methyl ethyl ketone or acetone or an alcohol such as methanol or ethanol. A mixture of dichloromethane and methanol gave especially good yields. The reducing agent may be present during the reaction or added after ozonide formation is completed. Thus, for example tetracyanoethylene may be present in solution in acetone during ozonolysis. While reducing agents such as dimethyl sulphide may be used to reduce the ozonide after its formation, preferred reagents are trivalent phosphorus compounds such as triphenylphosphine.

Where an alcohol solvent is used, the aldehyde product of formula I may form an acetal derivative with the alcohol. This may, however, readily be cleaved hydrolytically, for example using an aqueous base e.g. sodium bicarbonate. The reaction is preferably carried out at low temperatures, for example, -78°C.

After modification of the 17-side chain, the residue X may be removed to yield, as indicated above, a 5,6-trans vitamin of general formula IV. The removal of the residue X will be effected in different ways, depending on its nature.

Where X is SO<sub>2</sub>, it is conveniently removed by thermolysis under basic conditions, e.g. in the presence of a hydroxylic solvent such as an alcohol, e.g. ethanol, containing a base such as an alkali metal carbonate, e.g. sodium carbonate.

Where X is a group of formula V, removal can readily be effected by removal of the —CO—W—CO—moiety, for example by basic hydrolysis or treatment with hydrazine, followed by mild oxidation of the unsubstituted vitamin hydrazide so formed to the corresponding azo-compound which spontaneously decomposes to yield the required 5,6,-trans vitamin. Basic hydrolysis can be effected using strong alkali such as sodium or potassium hydroxide, for example in solution in an alcohol such as methanol, or by treatment with an amine such as triethylamine. The preferred method, however, is treatment with hydrazine which produces the desired hydrazide in high yield; this reaction has not previously been described for decomposition of such Diels-Alder adducts. Oxidation may be effected using reagents capable of oxidising hydrazo compounds to azo compounds, for example ceric, cupric, ferric, ferricyanic or periodate salts or air. A preferred mild reagent, however, is a diaryl telluroxide such as dianisyl telluroxide, preferably used with a reoxidant such as 1,2-dibromotetrachloroethane and a base such as K<sub>2</sub>CO<sub>3</sub> as described in our British Patent Specification No. 2058758A.

Where a 1α-hydroxy vitamin D compound is required, the modified 5,6-trans-vitamin compound of formula IV, which carries the desired 17-side chain, optionally protected, may be subjected to 1α-hydroxylation, using the procedure of our South African patent No. 79/5958. Thus, the 5,6-trans vitamin compound may be reacted with a selenite ester, preferably formed *in situ* by reaction of selenium dioxide and an alcohol such as methanol. The quantity of selenium compound may be reduced if a re-oxidant is employed, for example a periodate salt or N-methyl morpholine 1-oxide.

Alternatively, a reactive derivative of a 22-hydroxy derivative of formula I or IV above may be 1q-hydroxylated by the above procedure and the desired side-chain built up subsequently.

The 5,6-trans vitamin D compound of formula IV, after modifications such as those described above, may readily be isomerised in high yield to a required active cis-vitamin compound by known techniques, for example by irradiation in the presence of iodine or diphenyl selenide or, preferably, a triplet photosensitizer having a triplet energy of the order of 45±5 Kcal per mole, such as anthracene, acridine or phenazine. To avoid isomerization to undesired tachysterol derivatives, acid conditions should be avoided and the photoisomerisation is preferably effected in the presence of a base such as triethylamine.

Where protected hydroxyl groups are present in the vitamin product, these may be removed by conventional methods. In general, the vitamin structure is somewhat sensitive to acids, but is resistant to basic conditions and the latter are advantageously used. Acyloxy groups can thus be removed using alkali metal hydroxide in an alcohol solvent such as methanol. Silyl groups may be removed by treatment with quaternary ammonium fluorides such as tetra-n-butylammonium fluoride. Since most of the reactions described above can be applied to compounds having unprotected hydroxyl groups, protecting groups may be removed, if desired, at various stages. Although the vitamins are resistant to bases (and sensitive to acids), the dienophile adducts tend to be sensitive to bases and relatively resistant to acids. Consequently, acid conditions may be used to deprotect hydroxyl groups at stages where the dienophile residue X is present.

In general, most of the stages described ab ve proceed in excellent yield. Wh n conditions ar optimised, yields of the order of 80% or more at each stage have been achieved. This renders the ov rall yi ld of modified vitamin, starting from vitamin D<sub>2</sub>, markedly better than those achieved using many previously suggested routes.

Th following Examples are given by way of illustration only:-

Microanalyses and mass spectra were obtained by the staff at the Institut de Chemie des Substances Naturelles du CNRS, Gif-sur-Yvette, France. Melting points were determined using either a Kofler block, M I-temp or Fisher-Johns apparatus and are uncorrected. Optical rotations were measured at room temperature using a Rudolph Photoelectric Polarimeter, Model 70, and are r ported for chl roform solutions unless otherwise stated. UV spectra were recorded using a Carey 11 spectrophotometer and are reported for ethanol solutions. The molar extinction coefficient (ε) for these absorbances are given in parenthesis. IR spectra were recorded using a Perkin-Elmer 137 "Infracord" spectrophotometer and are reported for KBr discs unless otherwise stated. Absorbance characteristics are denoted by s = strong, m = medium, w = weak, sh = shoulder, br = broad. 'Hnmr spectra were determined at 60MHz on a Varian T-60 spectrometer. NMR characteristics are denoted as s = singlet, d = doublet, tr = triplet, q = quartet, m = multiplet, W = peak width at half height and are reported for CDCl<sub>3</sub> solutions, unless otherwise indicated, with tetramethylsilane as internal standard, as values of δ (ppm downfield of TMS).

Thin layer chromatography (tlc) was carried out on 250µ silica gel GHLF "Uniplates" (Analtech, USA); and preparative layer chromatography (plc) on 1 mm silica gel GF-254 "Uniplates" (Analtech, USA). "Chromatography" refers to medium pressure liquid chromatography carried out using E. Merck silica gel 60H. High performance liquid chromatography (HPLC) was carried out using Waters Associates silica gel "Porasil A" packed in two 2 ft × 3/8 inch stainless steel columns, and a Waters Associates chromatograph, equipped with a 6000 psi pump and a differential refractometer detector. Ozone was generated from a Towers Ozone Apparatus GE-150. Selective ozonolysis requires vigorous mixing of the dissolved substrate and the oxygen-ozone gaseous mixture. A "Vibromixer" (Chemapag, Switzerland) equipped with a stainless steel gas inlet/stirrer was particularly useful for this purpose. This equipment was also used for the formation of the phthalazine-1,4-dione Diels-Alder adducts of vitamin D.

A 200W Hanovia medium pressure mercury vapour lamp (654A36) was used as irradiation source fo 5,6-double bond photoisomerisation reactions.

Reactions on calciferol substrates were routinely performed under an inert, argon atmosphere. Calciferols were stored at  $-20^{\circ}$ C, under argon, in the dark, as either crystalline solids or (where possible) ether solutions. Solvents used were reagent grade unless otherwise stated.

Aqueous work-up refers to partition between an organic solvent and water, followed by sequential washing with a 5% aqueous sodium bicarbonate solution and a saturated aqueous sodium chloride solution. The organic solution was dried using either anhydrous MgSO<sub>4</sub> or anhydrous Na<sub>2</sub>SO<sub>4</sub>, and the solvent removed on a rotary evaporator. Acid work-up refers to partition between an organic solvent and water, followed by sequential washing with a 4% aqueous HCl solution; 5% aqueous sodium bicarbonate solution, etc. as for aqueous work-up.

Example 1

(a) 6(R),19-[4'-phenyl-1',2',4'-triazolidine-3',5'-dione-1',2'-yl]-9,10-sec-3β-hydroxy-ergosta-5(10), 7(E),

To ergocalciferol (5 g) in ethylacetate (150 ml) at 0°C under an argon atmosphere, 4-phenyl-1,2,4-triazoline-3,5-dione (2.4 g, 1.1 eq) in ethyl acetate (150 ml) was added over 45 min. After a further 1 hr, some of the title adduct had precipitated. The mixture was filtered and the filtrate passed down a neutral alumina column. Elution with hexane/ethylacetate gave the remainder of the product. Crystallisation from alcohol gave 6.2 g (86%). mp 99°C;  $[\alpha]_D = +208^\circ$  (c = 0.76);  $^1$ Hnmr  $\delta$  7.48 (s, 5H, aryl), 5.22 (m, W=10Hz, C—22H, 23H), 4.98 and 4.73 (an AB system, J=10Hz, C—6H, 7H), 4.2 and 3.85 (an AB system, J=15Hz, C—19H<sub>2</sub>), 4.1 (m, C—3H), 0.533 (s, C—18H<sub>3</sub>). IR vmax (CHCl<sub>3</sub>) 3700 (br), 2950 (s), 1775 (m), 1710 (s), 1425 (s)cm<sup>-1</sup>; mass spec. molecular ion, m/e = 571; (analysis found: % C, 75.63; H, 8.62; N, 7.36; C<sub>38</sub>H<sub>49</sub> $\mathcal{G}_3$ N<sub>2</sub>; requires: % C, 75.62; H, 8.64; N, 7.35).

Similarly prepared from ergocalciferol acetate in 85% yield was the corresponding acetate 6(R), 19-[4'-phenyl-1',2',4'-triazolidine-3',5'-dione-1',2'-yl]-9, 10-seco-3β-acetoxy-ergosta-5(10), 7(E), 22(E)-triene.

Crystallised from ethanol.m.p. 85°C;  $[\alpha]_D = +183^\circ$  (c = 0.82); <sup>1</sup>Hnmr  $\delta$  7.48 (s, 5H, aryl), 5.22 (m, W=12Hz, C—3H, 22H, 23H), 4.98 and 4.73 (an AB system, J=10Hz, C—6H, 7H), 4.2 and 3.85 (an AB system J=16Hz, C—19H<sub>2</sub>), 2.0 (s, OAc), 0.53 (s, C—18H<sub>3</sub>); IR vmax (CHCl<sub>3</sub>) 2950 (s), 2900 (sh), 1725 (s), 1420 (m)cm<sup>-1</sup>; mass spec. molecular ion m/e = 613; (analysis found: % C, 74.18; H, 8.11; N, 6.65;  $C_{38}H_{51}O_4N_3$ ; requires: % C, 74.35; H, 8.38; N, 6.85).

(b) Ozonolysis

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The adduct from (a) above (250 mg) in acetone (10 ml) containing tetracyanoethylene (55 mg, 1 eq) at -78°C was treated with ozone for 3 min (approx. 1.5 eq). The system was purged with argon whilst warming to room temperatur. The product mixture was separated by plc to give 130 mg of starting material (nmr) and the c rresponding 20(S)-f rmyl derivative (90 mg, 84%) as a white foam. ¹Hnmr δ 9.55 (d, J=3.75Hz, C—22H), 7.45 (s, 5H, aryl), 5.15 (m, W = 12Hz, C—3H), 4.92 and 4.82 (an AB system, J=10Hz, C—6H, 7H), 4.18 and 4.70 (an AB system J=16Hz, C—19H<sub>2</sub>), 2.0 (s, OAc), 1.12 (d, J=7H, C—21H<sub>3</sub>), 0.57 (s, C—18H<sub>3</sub>).

## Example 2

(a) Reaction of ergocalciferol acetate with phthalazine-1,4-dion

(b) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-hydroxy-ergosta-5(10),7(E), 22(E)-triene

To the acetate from (a) above (5 g) in benzene (100 ml) were added NaOH/CH<sub>3</sub>OH (1.25 M solution 12 ml). After 20 min, the mixture was diluted with water and  $CH_2Cl_2$ . Acid work-up gave an essentially quantitative yield (4.5 g) of the title 3 $\beta$ -hydroxy compound, crystalline from  $CH_2Cl_2$ /ether. m.p. 169—171°C; [ $\alpha$ ]<sub>D</sub> = +392°. (c=0.773); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.12 (m, W=9Hz, C—22H, 23H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.1 (m, C—3H), 0.18 (s, C—18H<sub>3</sub>). IR v max 3550 (br), 2950 (s), 2900 (sh), 1650 (s), 1610 (m), 1375 (m), 1350 (m)cm<sup>-1</sup>; mass spec. molecular ion m/e = 556; (analysis found: % C, 77.76; H, 8.78, N, 5.17;  $C_{95}H_{48}O_3N_2$  requires: % C, 77.66; H, 8.69; N, 5.03.

(c) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-tetrahydropyranyloxyergosta-5(10), 7(E), 22(E)-triene The alcohol from (b) above (4.5 g) in benzene (100 ml) was stirred overnight with dihydropyran (10 ml) and p-toluene sulphonic acid (10 mg). Aqueous work-up gave the title THP ether (204c) (5 g, 96%). Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/ether. m.p. 151—154°C; [α]<sub>D</sub> = +332° (c = 1.25); ¹Hnmr δ 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W = 10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.07 (m, W=9Hz, C—22H, 23H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.7 (m, THP, C—2'H), 4.02 (m, C—3H), 3.5 (m, W=20Hz, THP, C—6'H<sub>2</sub>), 0.17 (s, C—18H<sub>3</sub>); IR v max 2950 (s), 2900 (sh), 1650 (s), 1610 (m), 1370 (s), 1350 (s), 1030 (s)cm<sup>-1</sup>; mass spec. molecular ion m/e = 640.

(d) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-[t-butyldimethylsilyloxy]ergosta-5(10), 7(E), 22(E)-triene
 The alcohol from (b) above (4.5 g) in CH<sub>2</sub>Cl<sub>2</sub> (20 ml) was treated with t-butyl-dimethylsilylchloride (1.9 g) and imidazole (2.7 g) at room temperature for 1.5 hr. Addition of water followed by acid work-up and crystallization from CH<sub>2</sub>Cl<sub>2</sub>/hexane gave 5.1 g (94%) of the silyl ether. m.p. 203—205°C; [α]<sub>0</sub> = +313° (c = 1.5); ¹Hnmr δ 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.08 (m, W=9Hz, C—22H, 23H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.03 (m, C—3H), 0.88 (s, t-butyl), 0.17 (s, C—18H<sub>3</sub>), 0.07 (s, Si—CH<sub>3</sub>), 0.05 (S, Si—CH<sub>3</sub>); IR v max 2950 (s), 2900 (sh), 1650 (s), 1610 (m), 1370 (s), 1350 (s), 1090 (s)cm<sup>-1</sup>; mass spec. molecular ion m/e = 670; (analysis found: % C, 74.98; H, 9.26; N, 4.13; C<sub>42</sub>H<sub>62</sub>O<sub>3</sub>N<sub>2</sub>Si requires: % C, 75.18; H, 9.31; N, 4.18.

(e) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3 $\beta$ -methoxyethoxyethoxyergosta-5(10), 7(E), 22(E)-triene The alcohol from (b) above (4.5 g) in  $CH_2CI_2$  (100 ml) was stirred overnight at room temperature with methoxyethoxymethylchloride (8 ml) in the presence of diisopropylethylamine (20 ml). Acid work-up followed by chromatography and crystallisation from  $CH_2CI_2$ /hexane gave 4.3 g (83%) of the MEM ether. m.p. 123—125°C; [ $\alpha$ ]<sub>D</sub> = +325° (c = 1.295); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.15 (m, W=9Hz, C—22H, 23H), 4.82 (s, —OCH $_2$ O—), 4.78 and 4.22 (an AB system, J=18Hz, C—19H $_2$ ), 4.75 (d, J=10Hz, C—6H), 4.0 (m, C—3H), 3.67 (m, W=6Hz, —OCH $_2$ CH $_2$ O—), 3.43 (s, OCH $_3$ ), 0.18 (s, C—18H $_3$ ); IR  $\gamma$  max 2950 (s), 2900 (sh), 1650 (s), 1610 (m), 1470 (m), 1450 (m), 1370 (s), 1340 (s), 1100 (s), cm $^{-1}$ ; mass spec. molecular ion m/e = 644; (analysis found: % C, 74.72; H, 8.57; N, 4.13;  $C_{40}H_{56}O_5N_2$  requires: % C, 74.50; H, 8.75; N, 4.34.

(f) General procedure for the ozonolysis of the ergostane side chain

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The adduct (from (a), (c), (d) or (e) above (4—5 g) in CH₂Cl₂ (130 ml) and methanol (60 ml) was cooled to −78°C. The efficiently mixed solution was treated with an ozin -oxygen mixtur (approx. 1 mmol 0₃/min) for 8—12 min (tlc control) and then thoroughly purged with dry argon for approx. 5 min. Triphenylphosphine (2.5—3 g) was added and the mixture, after approx. 30 min at −78°C (tlc monitoring of the breakdown

of the methoxyhydroperoxide intermediates) was shaken with 5% aqueous NaHCO<sub>3</sub> (to prevent dimethyl acetal formation) and allowed to warm to room temperature. The layers were separated and the organic solution dried. Chromatography through silica gel (40—50 g) gave the aldehyde (75—86%) free from any of the C—20 (R) epimer (nmr). The following compounds were prepared in this manner.

1) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-acetoxy-20(S)-f rmylpregna-5(10),7(E)-diene

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Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/ether. m.p. 192—193°C;  $[\alpha]_D = +382$ °(c = 1.235); <sup>1</sup>Hnmr  $\delta$  9.55 (d, J=3Hz, C—22H), 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.17 (m, C—3H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 2.07 (s, OAc), 1.07 (d, J=7Hz, C—21H<sub>3</sub>), 0.22 (s, C—18H<sub>3</sub>); IR vmax (CHCl<sub>3</sub>) 2950 (m), 2900 (sh), 1740 (s), 1645 (s), 1610 (m), 1370 (m), 1350 (m), cm<sup>-1</sup>; mass spec. molecular ion m/e = 530; (analysis found: % C, 72.13; H, 7.12; N, 5.20; C<sub>32</sub>H<sub>38</sub>O<sub>5</sub>N<sub>2</sub> requires: % C, 72.43; H, 7.22; N, 5.28).

2) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-tetrahydropyranyloxy-20(S)-formyl-pregna-5(10), 7(E)-diene

Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/ether. m.p. 154—156°C;  $[\alpha]_D] = +356$ ° (c = 0.84); <sup>1</sup>Hnmr  $\delta$  9.42 (d, J=3Hz, C—22H), 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.69 (m, THP, C—2'H), 4.0 (m, C—3H), 3.5 (m, W=18Hz, THP, C—6'H<sub>2</sub>), 0.95 (d, J=6Hz, C—21H<sub>3</sub>), 0.23 (s, C—18H<sub>3</sub>). IR vmax 2950 (s), 2900 (sh), 1725 (s), 1640 (s), 1610 (m), 1370 (m), 1350 (m), 1025 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 572; (analysis found: % C, 72.89; H, 7.58; N, 4.78;  $C_{35}H_{44}O_5N_2$  requires: % C, 73.40; H, 7.74; N, 4.89).

3) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-[t-butyldimethylsilyloxy]-20(S)-formyl-pregna-5(10), 7(E)-

Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/hexane. m.p. 195—197°C;  $[\alpha]_D = +335^\circ$  (c = 1.64); <sup>1</sup>Hnmr  $\delta$  9.52 (d, J=3Hz, C—22H), 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.07 (m, C—3H), 1.07 (d, J=7Hz, C—21H<sub>3</sub>), 0.88 (s, t-butyl), 0.22 (s, C—18H<sub>3</sub>), 0.07 (s, Si—CH<sub>3</sub>), 0.03 (s, Si—CH<sub>3</sub>); IR vmax 2950 (s), 2900 (sh), 1740 (s), 1650 (s), 1610 (s), 1350 (s), 1090 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 602; (analysis found: % C, 71.57; H, 8.49; N, 4.51;  $C_{36}H_{50}O_4N_2Si$  requires: % C, 71.72; H, 8.36; N, 4.65).

4) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-20(S)-formyl-3β-methoxyethoxymethoxy-pregna-5(10), 7(E)-diene

Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/hexane, m.p. 136—137°C;  $[\alpha]_0 = +327^\circ$  (c = 0.62); <sup>1</sup>Hnmr  $\delta$  9.49 (d, J=3Hz, C—22H), 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 4.87 (s, —OCH<sub>2</sub>O—), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.03 (m, C—3H), 3.7 (m, W=6Hz, —OCH<sub>2</sub>CH<sub>2</sub>O—), 3.47 (s, OCH<sub>3</sub>), 1.07 (d, J=7Hz, C—21H<sub>3</sub>), 0.22 (s, C—18H<sub>3</sub>); IR vmax 2950 (m), 2900 (sh), 1740 (m), 1650 (s), 1610 (m), 1370 (m), 1350 (m), 1030 (m).

## Example 3

General procedure for the reduction of the C—20(S)-formyl to the C—20(S)-(hydroxyethyl) derivative The aldehyde compound (2.5—3.5 g) in benzene (60—90 ml) was added dropwise over a 15—20 min period to NaBH₄ (0.8—1.0 g) in ethanol (20—30 ml). After the addition, the excess reducing agent was carefully quenched with dilute aqueous HCl. The mixture was diluted with CH₂Cl₂. Aqueous work-up gave the desired alchohol in essentially quantitative yield. The following compounds have been prepared in this manner.

1) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-acetoxy-20(S)-[hydroxymethyl]-pregna-5(10), 7(E)-diene Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/ether. m.p. 238—240°C; [α]<sub>D</sub> = +363° (c = 0.875); <sup>1</sup>Hnmr δ 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.07 (m, C—3H), 4.78 and 4.21 (an AB system, J=18Hz, C—19H<sub>2</sub>); 4.75 (d, J=10Hz, C—6H), 3.47 (m, W=14Hz, C—22H<sub>2</sub>), 2.05 (s, OAc), 1.0 (broad singlet, C—21H<sub>3</sub>), 0.17 (s, C—18H<sub>3</sub>); IR vmax (CHCl<sub>3</sub>) 3200 (br), 2950 (m), 2900 (sh), 1750 (m), 1650 (s), 1610 (m), 1380 (m), 1350 (m), cm<sup>-1</sup>; mass spec. molecular ion m/e = 532.

2) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-20(S)-[hydroxymethyl]3β-tetrahydropyranyloxy-pregna-5(10), 7(E)-diene

Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/ether, m.p. 170—173°C;  $[\alpha]_D = +341^\circ$  (c = 0.58); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.67 (m, THP, C—2'H), 4.0 (m, C—3H), 3.5 (m, W=18Hz, C—22H<sub>2</sub>, THP, C—6'H<sub>2</sub>), 1.0 (broad singlet, C—21H<sub>3</sub>), 0.19 (s, C—18H<sub>3</sub>); IR vmax 3600 (br), 2950 (s), 2900 (sh), 1650 (s), 1610 (m), 1370 (m), 1350 (m), 1025 (m), cm<sup>-1</sup>; mass spec. molecular ion m/e = 574; (analysis found: % C 72.96; H, 7.96; N, 4.73; C<sub>35</sub>H<sub>46</sub>O<sub>5</sub>N<sub>2</sub> requires: % C, 73.14; H, 8.07; N, 4.87).

3) 6(R),19-[N,N'-phthalhydrazido]-9,10-sec -3β-[t-butyldimethylsilyloxy]-20(S)-[hydroxymethyl]-pr gna-5(10), 7(E)-diene

Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/hexane. m.p. 145—148°C;  $[a]_0 = +312^\circ$  (c = 1.22); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 4.78 and 4.22 (an AB syst m, J=18Hz, C—19H<sub>2</sub>),

4.75 (d, J=10Hz, C=6H), 4.03 (m, C=3H), 3.4 (m, W=14Hz,  $C=22H_2$ ), 1.0 (broad singlet,  $C=21H_3$ ), 0.88 (s, t-butyl), 0.19 (s,  $C=18H_3$ ), 0.07 (s,  $C=CH_3$ ); IR vmax 3500 (br), 2950 (s), 2900 (sh), 1640 (s), 1610 (m), 1340 (s), 1250 (s), 1090 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 604; (analysis found: % C, 71.56; H, 8.70; N, 4.47;  $C_{36}H_{52}O_4N_2S$ i requires: % C, 71.48; H, 8.67; N, 4.63).

#### Example 4

6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-acetoxy-20(S)-ethenylpregna-5(10), 7(E)-diene

Methyltriphenylphosphonium bromide (60 mg, 1.2 eq) was suspended in THF (6 ml). n-Butyl lithium (1.5 M solution 0.15 ml) was added. To the resulting orange-coloured solution, the 3β-acetoxy aldehyde from Example 2(f) (1) (100 mg) in benzene (6 ml) was added quickly. After a further 10 min, water was added and the mixture extracted with  $CH_2Cl_2$ . Acid work-up followed by purification by plc gave 75 mg (75%) of the title product. Crystalline from  $CH_2Cl_2$ /ether. m.p. 173—175°C; [α]<sub>D</sub> = +386° (c = 0.86); <sup>1</sup>Hnmr δ 8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.6—4.8 (m, C—3H, 22H, 23H<sub>2</sub>), 4.78 and 4.21 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 2.03 (s, OAc), 0.95 (d, J=7Hz, C—21H<sub>3</sub>), 0.17 (s, C—18H<sub>3</sub>); IR vmax 2950 (m), 1740 (s), 1650 (s), 1610 (m), 1370 (s), 1350 (s), 1260 (s), 1230 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 528; (analysis found: % C, 75.03; H, 7.72; N, 5.21;  $C_{33}H_{40}O_4N_2$  requires % C, 74.97; H, 7.63; N, 5.30).

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## Example 5

(a) Preparation of isobutylene epoxide

To methylallyl chloride (200 ml), 186 g) cooled in an ice bath was added 80%  $H_2SO_4$  ( $H_2SO_4$ , 95%, 109 ml;  $H_2O$ , 40 ml, 1 eq) over a 30 min period. The temperature of the mixture was maintained between 5—10°C. After a further 3 hr the mixture was added to ice and diluted to a total volume of approx. 100 ml. The layers were separated and the organic residue distilled to remove the by-product  $\beta$ ,  $\beta$ -dimethylvinyl chloride and unreacted starting material. These materials are removed below 80°C. The darkly coloured distillation residue is 1-chloro-2-methyl-propan-2-ol (128a)  $\delta$  3.47 (s, 2H), 2.97 (s, 1H, exchanges with  $D_2O$ ), 1.32 (s, 6H). This material was used without further purification.

To a 500 ml round bottom flask containing KOH (200 g) in water (125 ml) at 80°C and fitted with a mechanical stirrer and condenser, was added dropwise the crude chlorohydrin. The crude epoxide distilled directly from the reaction mixture. Redistillation gave isobutylene epoxide (48 g, 35%), b.p. 51°C (lit. 128°C 52°C); 14nmr δ 2.6 (s, 2H), 1.33 (s, 6H).

#### (b) 4-bromo-2-methyl-2-hydroxy-butane

To ethyl-3-bromo-propionate (21 g) in ether (150 ml) at 0°C was added methylmagnesium bromide (3 M soln. in ether, 125 ml, excess) dropwise. After the addition was complete, the mixture was stirred for a further 2 hrs at room temperature. After cooling again to 0°C, the mixture was carefully quenched with NH<sub>4</sub>Cl (30 g) in water (200 ml). The layers were separated and the ether layer washed with water until neutral, followed by brine, and dried. Evaporation gave the crude bromo-alcohol (227a)  $^1$ Hnmr  $\delta$  3.53 (t, J=9Hz, 2H), 2.93 (s, 1H, exchanges with D<sub>2</sub>O), 2.07 (t, J=9Hz, 2H), 1.27 (s, 6H); [lit.  $^{171}$  3.54 (t, J=8.5Hz, 2H), 2.65 (s, broad, 1H), 2.10 (t, J=8.5Hz, 2H), 1.26 (s, 6H)].

#### (c) 4-Bromo-2-methyl-2(triethylsilyloxy)-butane

Half of the crude bromide from (b) above in ether (50 ml) containing pyridine (5 ml), imidazole (10 g) and triethylsilylchloride (10 ml) was stirred for 2 days at room temperature. Water was added. Acid-work-up followed by chromatography gave 11 g (62% from the propionate) of the desired compound, homogeneous by tlc.  $^1$ Hnmr  $\delta$  3.52 (m, 2H), 2.03 (m, 2H), 1.25 (s, 6H), 1.2—0.2 (m, 15H); IR vmax (thin film) 3000 (s), 2950 (sh), 1460 (m), 1420 (m), 1380 (m), 1365 (m), 1230 (s), 1195 (s), 1170 (m), 1100 (s), 1040 (s), 1010 (s), 965 (m), 840 (w), 740 (s), 720 (s), cm $^{-1}$ .

#### (d) 3-Methyl-2-buten-1-yl-triphenylphosphonium bromide

The bromide from (c) above (1 g) and triphenylphosphine (0.9 g) in benzene (4 ml) was thoroughly degassed, and then heated to reflux. After 3 days, the insoluble material was filtered off to give 1.2 g (85%) of phosphonium salt (228). Recrystallised from CH<sub>2</sub>Cl<sub>2</sub>/ETOAc. m.p. 234—238°C (lit.<sup>119b</sup> 236—239°C); <sup>1</sup>Hnmr  $\delta$  8.17—7.67 (m, 15H, aryl), 5.18 (m, W=18Hz), 4.73—4.2 (m, 2H), 1.67 (d, J=5Hz, 3H), 1.31 :d, J=5Hz, 3H); IR vmax 2900 (w), 1590 (w), 1490 (m), 1435 (s), 1110 (s), cm<sup>-1</sup>.

## (e) Methyldiphenylphosphine oxide

Methyltriphenylphosphonium bromide (6 g) was refluxed overnight with KOH (5 g) in water (70 ml). The mixture was allowed to coll to room temperatur and the normal xtracted (3x) with  $CH_2Cl_2$ . The organic layer was washed with brine, dried and the solvent removed to give the crude solid product (3.5 g) in essintially quantitative yield. Ricrystallised from acetone. m.p. 113—114°C (lit. 132 109—111°C); 1Hnmr  $\delta$  8.0—7.3 (m, 10H, aryl), 2.03 (d, J=13Hz, 3H); IR vmax 1440 (s), 1175 (s), cm<sup>-1</sup>; mass spec. molecular ion m/ = 216.

(f) 3-Hydroxy-3-methylbut-1-yl-diphenylphosphine oxide

Methyldiphenylphosphine oxide (1.5 g) was suspended in ether (20 ml) at 0°C. BuLi (1.2 eq) was slowly added, and an orange coloured solution formed. To this was added isobutylene epoxide (0.8 ml, 1.3 eq). After approx. 15 min, the mixture was carefully quenched with water. This mixture was extracted with 5 CH<sub>2</sub>Cl<sub>2</sub> (2x) and the organic layer was washed with 4% aqueous HCl/brine and concentrated. The resulting yellow, oily product was dissolved in a water-ether mixture and the layers separated. The ether lay represented once with water, and the combined aqueous fractions extracted with CH<sub>2</sub>Cl<sub>2</sub> (3x). The organic layer was washed with brine and dried. The solvent was removed and the resulting colourless oil was taken up in benzene and refluxed through a soxhlet containing CaH<sub>2</sub> for 2 hr. The solvent was removed to give crude title compound (1.4 g, 70%; as an oil. ¹Hnmr δ 8.0—7.3 (m, 10H, aryl), 2.5 (m, W=34Hz, 2H), 1.83 (m, W=32Hz, 3H), 1.23 (s, 6H); IR vmax (CCl<sub>4</sub>) 3500 (m), 2950 (m), 1440 (s), cm<sup>-1</sup>.

(g) 3-tetrahydropyranyloxy-3-methylbut-1-yl-diphenylphosphine oxide

The phosphine oxide from (f) above (1.4 g) was dissolved in dihydropyran (20 ml) and benzene (5 ml).

p-Toluenesulphonic acid (10 mg) was added. After 20 hr, the mixture was concentrated, added to CH<sub>2</sub>Cl<sub>2</sub> and washed with 5% aqueous NaHCO<sub>3</sub>/brine and dried. Evaporation of the solvent gave the crude product (1.8 g) essentially quantitatively as a solid. Recrystallised from acetone. m.p. 146—148°C; ¹Hnmr δ 8.0—7.3 (m, 10H, aryl), 4.67 (m, w=6Hz, THP, C—2'H), 3.67 (m, W=36Hz, THP, C—6'H<sub>2</sub>), 1.23 (s, 6H); IR vmax 2950 (m), 1440 (m), 118C (s), cm<sup>-1</sup>; (analysis found: % C, 70.80; H, 7.73; P, 8.54; C<sub>22</sub>H<sub>29</sub>O<sub>3</sub>P requires % C, 70.96; H, 7.85; P, 8.32.

(h) Preparation of the lithium bromide adduct of the betaine

To methyltriphenylphosphonium bromide (2.898 g) suspended in ether (50 ml) cooled to 0°C was added butyl lithiumn (2.03 M soln.; 4 ml). Isobutylene epoxide (1.0 ml, 1.25 eq) was added and some insoluble material instantly formed. After stirring for 15 min, the reaction mixture was allowed to settle and the supernatant liquid was removed. The resulting solid was suspended in ether and transferred to two centrifuge tubes, and spun. The ether was removed. This process was repeated until the ether washing were colourless (usually 4x). The colourless solid material was dried to give the Li-Br adduct of the betaine (1.5 g) 42%. Beilstein and lithium ion positive, flame tests. IR vmax (nujol) 3500 (br), 3000 (s), 1440 (s), cm<sup>-1</sup>.

(i) [3-(triethylsilyloxy)-3-methylbut-1-yl]-triphenylphosphonium tetraphenyl borate

To methyltriphenylphosphonium bromide (3 g) suspended in THF (40 ml) was added phenyl lithium (1 eq; 6 ml of a 1.5 M soln.). After 15 min isobutylene epoxide (1 ml, 1.25 eq) was added followed, after a further 5 min, by a second addition of phenyl lithium (1 eq). To this mixture was added benzophenone (1 g; approx. 0.3 eq). After stirring for 20 min, the reaction was quenched with 48% aqueous HBr until just acidic (litmus paper). The organic solvent was removed on a rotary evaporator, water was added and the aqueous layer washed with ether, and the layers separated. The water was removed (rotary evaporator) and the resulting oil taken up in CH<sub>2</sub>Cl<sub>2</sub>. Aqueous work-up gave the phosphonium salt (226) (3.1 g) 58% as an oil. <sup>1</sup>Hnmr δ 8.17—7.67 (m, 15H, aryl), 5.37 (broad s, —OH), 3.8 (m, W=32Hz C—1H<sub>2</sub>), 1.8 (m, W=22Hz, C—2H<sub>2</sub>), 1.28 (s, (—CH<sub>3</sub>)<sub>2</sub>); IR vmax (CHCl<sub>3</sub>) 3450 (s), 3000 (s), 1590 (sh), 1440 (s), cm<sup>-1</sup>.

(j) Silylation

To the phosphonium salt from (i) above (3.7 g) in  $CH_2Cl_2$  (70 ml) was added imidazole (3.4 g) followed by triethylsilylchloride (5 ml). After 40 hr stirring at room temperature, water was added and the mixture diluted with  $CH_2Cl_2$ . The  $CH_2Cl_2$  solution after an acid work-up was evaporated and the oily residue partitioned between water and hexane/ether. The water was evaporated and the residue taken up in  $CH_2Cl_2$  which was washed with brine and dried to give on evaporation the salt (226b) (3.6 g, 77%) as an oil.

(k) Anion exchange

To the phosphonium salt from (j) above (3.6 gg) in 95% ethanol (50 ml) was added dropwise, with stirring, a solution of sodium tetraphenyl borate (2.5 g; 1.1 eq) in water (20 ml). An oily residue is formed which solidifies on continued stirring. Filtration gives the phosphonium tetraphenyl borate salt (4.78 g, 92%) as a white, amorphous, non-hygroscopic solid which may be recrystallised from acetone/hexane/ethanol. m.p. 150—151°C;  $^1$ Hnmr  $\delta$  (acetone-d<sub>e</sub>) 8.2—6.8 (m, 35H, aryl), 3.53 (m, W=34Hz, C—1H<sub>2</sub>), 1.8 (m, W=24Hz, C—2H<sub>2</sub>), 1.33 (s, (—CH<sub>3</sub>)<sub>2</sub>), 1.25—0.5 (m, 15H, —SiEt<sub>3</sub>); IR vmax 3100 (s), 2950 (s), 1580 (m), 1490 (s), 1110 (s), 1020 (s), cm<sup>-1</sup>; (analysis found: % C, 81.41; H, 7.73; P, 3.93; C<sub>53</sub>H<sub>60</sub>BOPSi requires: % C, 81.31; H, 7.73; P, 3.96.

(I) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-acetoxy-25-hydroxy-cholesta-5(10),7(E),22(E)-triene

To methyltriph nylphosphonium bromide (2.898 g) suspended in THF (32 ml) at 0°C was added butyl lithium (2.03 M, 4 ml). Iso-butyl ne ep xide (720 µl, 1 eq) was slowly added. After a further 15 min, butyl lithium (4 ml) was added. To 3 ml of this solution was added the aldehyde from Example 2(f) (1) (300 mg) in benzene (10 ml). The red colour was quickly discharged. Water was added and the mixture extracted with 65 CH<sub>2</sub>Cl<sub>2</sub>. After acid work-up the major product was isolated by plc to give the title compound (105 mg, 31%).

#### Method B

Th b taine from (h) abov (628 mg) was suspended in ether (15 ml) and THF (10 ml). Butyl lithium was added dropwise until a stable colour was formed and then a further amount (0.75 ml, 2 eq for steroid, 1 eq for P compound) was added. To this mixture was added the aldehyde from Example 2(f) (1) (400 mg) in benzene (6 ml) (approx. 5 min). After the addition, water was added and the mixture xtracted with CH<sub>2</sub>Cl<sub>2</sub>. Work-up as above gave the title compound (155 mg, 34%).

#### Method C

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The phosphonium salt from (h) above (280 mg, 1.5 eq) was dissolved in THF (15 ml) at 0°C. Phenyl lithium (3 eq) was added. The aldehyde (206a) (150 mg, 1 eq) in benzene (6 ml) was added quickly. Tlc showed no change during 30 min and so water was added. Work-up as above, and isolation by plc gave th title product (80 mg, 47%). Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/ether. m.p. 175—177°C; [ $\alpha$ ]<sub>p</sub> = +347° (c = 0.83); ¹Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.27 (m, W=10Hz, C—3H, 22H, 23H), 44.78 and 4.21 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 2.03 (s, OAc), 1.15 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 0.97 (d, J=7Hz, C—21H<sub>3</sub>), 0.17 (s, C—18H<sub>3</sub>); IR vmax 3800 (m), 2950 (s), 2900 (sh), 1750 (s), 1650 (s), 1610 (m), 1370 (s), 1350 (s), 1240 (s), 965 (m), cm<sup>-1</sup>; mass spec. molecular ion m/e = 600; (analysis found: % C, 73.94; H, 8.17; N, 4.59; C<sub>37</sub>H<sub>48</sub>O<sub>5</sub>N<sub>2</sub> requires: % C, 73.97; H, 8.05; N, 4.66).

(m) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β-acetoxy-25-hydroxy-cholesta-5(10),7(E),22(Z)-triene

To the phosphonium salt from (i) above (1.9 g) in THF (30 ml) was added phenyl lithium (1.5 M soln., 1.7 ml, 1 eq). After a few minutes, the aldehyde (206a) (1 g) in benzene (35 ml) was added dropwise over about 1 min. After a further 3 min, water was added and the mixture diluted with CH₂Cl₂ and given an acid work-up. The reaction was repeated as above and the combined products chromatographed to yield 2.12 g (78%) of a crude, yellow coloured product.

The above mixture (1.4 g) was treated with AcOH: $H_2O$ :THF (8:1:1) (10 ml) for 1.5 hr. Dilution with CH<sub>2</sub>Cl<sub>2</sub> followed by aqueous work-up, chromatography and crystallisation gave 1 g of product (85%). Further recrystallisation from CH<sub>2</sub>Cl<sub>2</sub>/ether, indicated the major component to have the following characteristics. m.p. 182—184°C; [ $\alpha$ ]<sub>D</sub> = +339° (c = 0.84); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.27 (m, W=12Hz, C—3H, 22H, 23H), 4.78 and 4.21 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 2.03 (s, OAc), 1.17 (s,s C—26H<sub>3</sub>, 27H<sub>3</sub>), 0.9 (d, J=7Hz, C—21H<sub>3</sub>), 0.17 (s, C—18H<sub>3</sub>); IR vmax 3650 (m), 2950 (s), 2900 (sh), 1750 (s), 1650 (s), 1610 (m), 1370 (s), 1350 (s), 1240 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 600; (analysis found: % C, 74.10; H, 8.15; N, 4.47; C<sub>32</sub>H<sub>48</sub>O<sub>5</sub>N<sub>2</sub> requires: % C, 73.97; H, 8.05; N, 4.66).

(n) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3β,25-dihydroxy-cholesta-5(10),7(E)-diene

The unsaturated side chain compound from (i) above (450 mg) in benzene (5 ml) and ethanol (5 ml) containing NaHCO<sub>3</sub> (100 mg) and 5% Pt/C (150 mg) was stirred under a hydrogen atmosphere for 24 hr. The mixture was filtered through celite and the solvent removed. To the residue, in benzene (10 ml), was added NaOH in methanol (1.25 M soln, 2 ml) and the mixture stirred for 20 min at room temperature. Acid work-up and crystallisation from CH<sub>2</sub>Cl<sub>2</sub>/ether afforded 380 mg (91%) of the title side chain saturated diol. m.p. 174—177°C; [ $\alpha$ ]<sub>D</sub> = +408° (c = 0.825); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 4.78 and 4.22 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 4.11 (m, C—3H), 1.22 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 0.87 (broad singlet, C—21H<sub>3</sub>), 0.18 (s, C—18H<sub>3</sub>); IR vmax 3550 (s), 2950 (s), 2900 (sh), 1650 (s), 1610 (m), 1370 (s), 1350 (s), cm<sup>-1</sup>; (analysis found: % C, 74.65; H, 8.66; N, 5.06; C<sub>35</sub>H<sub>48</sub>O<sub>4</sub>N<sub>2</sub> requires: % C, 74.96; H, 8.63; N, 5.00).

## Example 6

General procedure for the conversion of the phthalazine-1,4-dione adduct to the corresponding 5(E),7(E),10(19)-triene system of the calciferol

The adduct (200—600 mg) was refluxed overnight, under argon in ethanol (10 ml) and hydrazine (3 ml). After cooling to room temperature, the solvents were removed under reduced pressure and the resulting solid taken up in water (30 ml) and CH₂Cl₂ (30 ml). To this two-phase system under argon was added dianisyltellurium oxide (150—450 mg), K₂CO₃ (6 g) and 1,2-dibromotetrachloroethane (3 g), and the mixtur stirred for approx. 5 hr (tic control). After acid work-up the mixture was chromatographed through silica gel (12 g) and the product removed from traces of tellurium oxidant by plc to give the desired vitamin D compound in 85—93% yield.

(1) 9,10-seco-3β,25-dihydroxy-cholesta-5(E),7(E),10(19)-triene

(2) 3B-(3',5'-dinitrobenzoate) ester

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The above crud calciferol (1) (125 mg) in pyridine (5 ml) was treated with 3,5-dinitrobenzoyl chlorid (85 mg, 1.1 eq). Water was added and the mixtur diluted with ther. Aft r acid work-up, the ster was is lated by plc (129 mg, 70%). Crystalline from ether/h xane, m.p. 105—107°C:  $\{\alpha\}_D = +168$  (c = 0.97); <sup>1</sup>Hnmr  $\delta$  9.13 (m, 3H, aryl), 6.62 and 5.82 (ABq, J=11Hz, C—6H, 7H), 5.3 (m, W=14Hz, C—3H), 5.07 (s, C—19H), 4.77 (s, C—19H), 1.23 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 0.93 (broad singlet, C—21H<sub>3</sub>), 0.43 (s, C—18H<sub>3</sub>); IR vmax 3550 (m), 2950 (s), 2900 (sh), 1750 (s), 1640 (w), 1550 (s), 1350 (s), 1275 (s), cm<sup>-1</sup>; (analysis found: % C, 68.62; H, 7.85; N, 4.65;  $C_{34}H_{46}N_2O_7$  requires: % C, 68.66; H, 7.80; N, 4.71).

#### Example 7

(a) 9,10-seco-3β,25-dihydroxy-cholesta-5(Z),7(E),10(19)-triene

A solution of the 5,6-trans compound from Example 6(1) (126 mg) in benzene (30 ml) containing triethylamine (2 drops) and anthracene (25 mg) was thoroughly degassed. A hanovia lamp (number 654A36) was placed such that the outside of the water cooled jacket was 15 cm from the reaction vessel. The mixture was irradiated for 25 min and the title 5,6-cis compound isolated by plc (93 mg, 74%). Crystalline from acetone/water. m.p. 98—100°C (lit.<sup>174</sup> 95—100°C); [ $\alpha$ ]<sub>D</sub> = +77° (c = 0.26); UV  $\lambda$ max 262 nm (19060); <sup>1</sup>Hnmr  $\delta$  6.25 and 6.1 (ABq, J=11Hz, C—6H, 7H), 5.05 (s, C—19H), 4.83 (s, C—19H), 3.9 (m, W=18Hz, C—3H), 1.27 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 0.95 (broad singlet, C—21H<sub>3</sub>), 0.55 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 2900 (sh), 1640 (w), 1480 (m), 1380 (m), 1055 (s), cm<sup>-1</sup>; (analysis: C<sub>27</sub>H<sub>44</sub>O<sub>2</sub> · H<sub>2</sub>O requires: % C, 77.46; H; 11.08; found: % C, 77.29; H, 11.08). The melting point of an authentic sample supplied by Roussel Uclaf, Inc. (Romainville France) did not depress on mixing.

(b) 3-(3',5-dinitrobenzoate) ester

Prepared as previously described in Example 6(2). Crystalline from ether/hexane. m.p. 149—150°C (lit.<sup>172</sup> 147—148°C);  $[\alpha]_D = +90^\circ$  (c = 0.6); (analysis:  $C_{34}H_{46}N_2O_7$  requires: % C, 68.66; H, 7.80; N, 4.71; found: % C, 68.94; H, 7.80; N, 4.52).

## Example 8

SO<sub>2</sub> adducts from 9,10-seco-3β-hydroxy-ergosta-5([Z),7(E),10(19),22(E)-tetraene

Sulphur dioxide was slowly passed through a well-stirred mixture of benzene (100 ml) and water (50 ml) containing ergocalciferol (5 g), for a total of 3.5 hr. After this time, air was passed through the mixture for approx. 20 min. Ether and brine were added and the layers separated. Aqueous work-up gave the known sulphur dioxide adducts (172a, 173a) which were used without further purification.

## Example 9

(a) 9,10-seco-3β-(triethylsilyloxy)-ergosta-5(E),777(E),10(19),22(E)-tetra-ene

To the 3 $\beta$ -alcohol corresponding to the title compound (4.3 g) in CH<sub>2</sub>Cl<sub>2</sub> (50 ml) was added imidazole (4 g) followed by triethylsilylchloride (3 ml). After a few minutes, water was added and the organic layer washed with water/brine and dried. The required silyl ether was isolated essentially quantitatively after chromatography as an oil. UV  $\lambda$ max 274 nm; <sup>1</sup>Hnmr  $\delta$  6.45 and 5.87 (ABq, J=11Hz, C—6H, 7H), 5.2 (m, W=9Hz, C—22H, 23H), 4.92 (s, C—19H), 4.63 (s, C—19H), 3.82 (m, W=18Hz, C—3H).

(b) 9,10-seco-1α-hydroxy-3β-(triethylsilyloxy)-ergosta-5(E),7(E),10(19),22(E)-tetraene

N-Methylmorpholine N-oxide (NMO) (6.3 g) was stirred with anhydrous MgSO<sub>4</sub> in CH<sub>2</sub>Cl<sub>2</sub> (50 ml) for 30 min. Selenium dioxide (1.3 g) was stirred in methanol (50 ml) for 45 min and warmed to reflux. The above CH<sub>2</sub>Cl<sub>2</sub> mixture was filtered into a solution of the 5,6-*trans*-ergocalciferol derivative from (a) above (5.5 g) in 1,2-dichloroethane (50 ml). This mixture was warmed to reflux and then the hot methanol mixture added, and refluxing of the whole continued for a further 35 min. The heat source was removed and the mixture diluted with CH<sub>2</sub>Cl<sub>2</sub>. Aqueous work-up followed by chromatography through silica gel (40 g) gave 2.66 g (47%) of the title compound as an oily product. UV λmax 274 nm; <sup>1</sup>Hnmr δ 6.57 and 5.90 (ABq, J=11Hz, C—6H, 7H), 5.25 (m, W=9Hz, C—22, 23H), 5.08 (s, C—19H), 4.98 (s, C—19H), 4.65—3.92 (m, C—1H, 3H).

(c) 9,10-seco-1α,3β-dihydroxy-ergosta-5(E(,7(E),10(19),22(E)-tetra-ene

The silylether from (b) above (460 mg) in THF (10 ml) was stirred for 30 min with tetrabutylammonium fluoride (460 mg). The mixture was diluted with  $CH_2CI_2$  and after aqueous work-up, the title diol was purified by plc to give 305 mg (84%). Crystalline from ether/hexane. m.p. 103—105°C; [ $\alpha$ ]<sub>D</sub> = +172° (c = 0.58); UV  $\lambda$ max 272 nm (22600); <sup>1</sup>Hnmr  $\delta$  6.38 and 5.82 (ABq, J=11Hz, C—6H, 7H), 5.18 (m, W=9Hz, C—22H, 23H), 4.9 (m, W=9Hz, C—19H<sub>2</sub>), 4.53—3.77 (m, C—1H, 3H), 0.57 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 2900 (sh), 1640 (w), 1460 (m), 1375 (m), 1050 (s), 1030 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 412; (analysis found: % C, 79.57; H, 10.71;  $C_{28}H_{44}O_2$  requires: % C, 81.50; H, 10.79;  $C_{28}H_{44}O_2$ .  $\frac{1}{2}H_2O$  requires: % C, 79.76; H, 10.76).

(d) 9,10-s co-1α-hydr xy-3β-triethylsilyloxy-erg sta-5(Z),7(E),10(19),22(E)-t tra- ne
The 5,6-trans compound from (b) above (600 mg) in benzene (30 ml) containing phenazin (120 mg) and triethylamine (few drops) was photoisomerised as ab ve for 30 min to give 400 mg (66%) of th

5,6-cis vitamin. UV  $\lambda$ max 263 nm;  $^1$ Hnmr  $\delta$  6.38 and 6.08 (ABq, J=11Hz, C—6H, 7H), 5.23 (m, W=10Hz, C—19H, 22H, 23H), 5.0 (s, C—19H), '4.6—3.92 (m, C—1H, 3H).

( ) 9,10-seco-1α-3β-dihydroxy-ergosta-5(Z),7(E),10(19),22(E)-tetra-ene

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The silyl ether derivative from (d) above (200 mg) was stirred at room temperature in THF (10 ml) with N-Bu<sub>4</sub>NF (1 M soln. in THF, 2 ml) for about 30 min. Dilution with CH<sub>2</sub>Cl<sub>2</sub> and aqueous work-up followed by purification by plc gave 129 mg (82%). Crystalline from ether/hexane gave the title compound. m.p. 141—143°C (lit.  $^{106}$  138—140°C); [ $\alpha$ ]<sub>D</sub> = +34° (c = 0.645); UV  $\lambda$ max 264 nm (19100);  $^{1}$ Hnmr  $\delta$  6.35 and 6.05 (ABq, J=11Hz, C—6H, 7H), 5.16 (m, W=14Hz, C—19H, 22H, 23H), 4.98 (s, C—19H), 4.6—3.85 (m, C—1H, 3H), 0.55 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 2900 (sh), 1640 (w), 1460 (m), 1370 (m), 1060 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 412; (analysis: C<sub>28</sub>H<sub>44</sub>O<sub>2</sub> requires: % C, 81.50; H, 10.75; O, 7.76; found: % C, 81.39; H, 10.60).

#### Example 10

(a) 9,10-seco-3β-acetoxy-1α-benzoyloxy-ergosta-5(E),7(E),10(19),22(E)-tetra-ene

The  $1\alpha$ -hydroxy-3 $\beta$ -triethylsilyloxy-compound from Example 9(b) (2 g) was treated with benzoyl chloride (2 ml) in pyridine (25 ml). After 30 min water was added and the mixture diluted with ether. After acid work-up, the solvent was removed and the resulting oil stirred overnight in THF: $H_2O$ :AcOH; 8:1:3 (36 ml). After dilution with ether and aqueous work-up, the crude benzoate-alcohol was taken up in benzene (40 ml). Triethylamine (7 ml), acetic anhydride (3 ml) and 4-dimethylaminopyridine (15 mg) were added. After 30 min, water was added and the mixture diluted with ether. Acid work-up and chromatography through silica (10 g) gave 1.76 g (83%) of the title acetate-benzoate as an oil.  $^1$ Hnmr  $\delta$  8.05 (m, W=12Hz, 2H, aryl), 7.5 (m, W=10Hz, 3H, aryl), 6.58 (d, J=11Hz, C—6H), 5.88 (m, W=16Hz, C—1H, 7H), 5.15 (m, W=10Hz, C—3H, 19H<sub>2</sub>, 22H, 23H), 2.05 (s, OAc), 0.57 (s, C—18H<sub>3</sub>).

(b) 6(R),19-[N,N'-phthalhydrazido]-9,10-seco-1α-benzoyloxy-3β-acetoxy-ergosta-5(10),7(E),22(E)-triene

To a well-stirred suspension of phthalhydrazide (2 g) in  $CH_2Cl_2$  (200 ml) at 0°C, containing the vitamin from (a) above (2 g), was added dropwise a solution of  $Pb(OAc)_4$  (4 g) in  $CH_2Cl_2$  (20 ml) and acetic acid (1 ml). After consumption of starting material (tlc control), the excess phthalhydrazide was removed by filtration. Aqueous work-up and chromatography gave 1.4 g (52% from 248b) of a 95:5 mixture of (250) and the presumed 6(S) isomer. Crystallisation from  $CH_2Cl_2/h$ exane gave pure title compound. m.p. 211—213°C;  $[cl_0]_0 = +295^\circ$  (c = 0.83);  ${}^1H$ nmr  $\delta$  8.5—7.3 (m, 9H, aryl), 5.95 (m, W=14Hz, C—1H, 7H), 5.28 (m, C—3H), 5.17 (m, W=10Hz, C—22H, 23H), 4.92 and 4.37 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.8 (m, C—6H), 2.05 (s, OAc), 0.17 (s, C—18H<sub>3</sub>): IR vmax 2950 (s), 2900 (sh), 1750 (s), 1720 (s), 1640 (s), 1610 (m), 1265 (s), 1245 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 718; (analysis found % C, 75.26; H, 7.54; N, 3.82;  $C_{45}H_{54}O_6N_2$ ; requires: %C 75.18; H, 7.57; N, 3.90).

#### Example 11

(a) SO<sub>2</sub> adducts of 9,10-seco-3β-(t-butyldimethylsilyloxy)-ergosta-5(E),7(E),10(19),22(E)-tetraene

The crude mixture of sulphur dioxide adducts of ergocalciferol (prepared from 5 g of ergocalciferol as described previously), in  $CH_2Cl_2$  (40 ml), containing imidazole (4 g) was stirred with t-butyldimethylsilyl chloride (3.5 g). After 1.5 hr, the reaction was worked-up as described previously to give, after chromatography, 4.8 g (66%, from ergocalciferol) of the title compound as an oil epimeric at C—6. <sup>1</sup>Hnmr  $\delta$  5.22 (m, W=9Hz, C—22H, 23H), 4.64 (m, W=10Hz, C—6H, 7H), 4.02 (m, W=16Hz, C—3H), 3.67 (broad s, C—19H<sub>2</sub>), 0.91 (s, t-Bu), 0.68 + 0.59 (2 × s, C—18H<sub>3</sub>), 0.07 s, [(Si—CH<sub>3</sub>)<sub>2</sub>].

(b) SO<sub>2</sub> adducts of 9,10-seco-3β-triethylsilyloxyl-ergosta-5(E),7(E),10(19),22(E)-tetra-ene

The crude mixture of sulphur dioxide adducts of ergocalciferol (prepared from 5 g of ergocalciferol as described previously), in  $CH_2Cl_2$  (40 ml), containing imidazole (4 g) was stirred with triethylsilylchloride (3.5 ml). After about 30 min, the reaction was worked up as described previously to give, after chromatography, 5.3 g (74% from ergocalciferol) of (210b) as an oil. <sup>1</sup>Hnmr  $\delta$  5.22 (m, W=9Hz, C—22H, 23H), 4.64 (m, W=10Hz, C—6H, 7H), 4.02 (m, W=16Hz, C—3H), 3.67 (broad s, C—19H<sub>2</sub>).

- - (d) Similarly pr pared in 82% yield from (b) above was the SO<sub>2</sub> adducts of 9,10-seco-3 $\beta$ -(triethylsilyl xy)-20(S)-formyl-pregna-5(E),7(E),10(19)-trien . <sup>1</sup>Hnmr  $\delta$  9.57 (m, C—22H), 4.67 (m, W=12Hz, C—6H, 7H), 3.97 (m, W=16Hz, C—3H), 3.65 (broad s, C—19H<sub>2</sub>), 1.15 (d, J=6Hz, C—21H<sub>3</sub>); IR vmax (thin film) 2950 (s), 2900 (sh), 1735 (s), 1660 (w), 1460 (m), 1380 (m), 1310 (s), 1150 (m), cm<sup>-1</sup>.

## Example 12

(a) SO<sub>2</sub> adducts of 9,10-seco-3 $\beta$ -(t-butyldimethylsilyloxy)-20(S)-(hydroxymethyl)-pregna-5(E),7(E),10(19)-triene

The aldehyd corresponding to the title compound (3.1 g) was reduced as described in the general procedure to the title compound in essentially quantitative yield,  $^1$ Hnmr  $^5$  4.63 (m, W=12Hz, C—6H, 7H), 4.02 (m, W=16Hz, C—3H), 3.80—3.28 (m, C—19H<sub>2</sub>, 22H<sub>2</sub>), 1.05 (d, J=6Hz, C—21H<sub>3</sub>), 0.87 (s, t-Bu), 0.68 + 0.58 ([2 × s, C—18H<sub>3</sub>), 0.05 [s, (Si—CH<sub>3</sub>)<sub>2</sub>]; IR max (thin film) 3550 (br), 2950 (s), 2900 (sh), 1660 (w), 1475 (m), 1350 (s), 1275 (s), 1155 (m), cm<sup>-1</sup>.

- (1) Similarly prepared in greater than 90% yield was the SO<sub>2</sub> adducts of 9,10-seco-3β-(triethylsilyloxy)-20(S)-(hydroxymethyl)-pregna-5(E),7(E),10(19)-triene, <sup>1</sup>Hnmr δ 4.63 (m, W=12Hz, C—6H, 7H), 3.93 (m, W=16Hz, C—3H), 3.77—3.17 (m, C—19H<sub>2</sub>, 22H<sub>2</sub>); IR vmax (thin film) 3550 (br), 2950 (s), 2900 (sh), 1660 (w), 1460 (m), 1380 (m), 1305 (s), 1240 (m), 1155 (m), cm<sup>-1</sup>.
- (2) 9,10-seco-3β-(t-butyldimethylsilyloxy)-20(S)-(hydroxymethyl)-pregna-5(E),7(E),10(19)-triene Adducts of (1) above (3 g) was stirred in refluxing methanol (50 ml) containing NaHCO<sub>3</sub> (3 g) for 2.5 hr. Work-up as described above gave 2.36 g (90%) of the calciferol. UV λmax 274 nm; ¹Hnmr δ 6.47 and 5.87 (ABq, J=11Hz, C—6H, 7H), 4.92 (s, C—19H), 4.65 (s, C—19H), 4.1—3.15 (m, C—3H, 22H<sub>2</sub>), 1.06 (d, J=5Hz, C—21H<sub>3</sub>), 0.9 (s, t-Bu), 0.58 (s, C—18H<sub>3</sub>), 0.07 s, [(Si—CH<sub>3</sub>)<sub>2</sub>].
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  (3) Similarly prepared in 47% yield from the adducts of (1) above after chromatography was 9,10-seco-3β-(triethylsilyloxy)-2;(S)-(hydroxymethyl)-pregna-5(E),7(E),10(19)-triene (267d). UV λmax 273 nm; ¹Hnmr δ 6.43 and 5.7 (ABq, J=11Hz, C—6H, 7H), 4.9 (s, C—19H), 4.6 (s, C—19H), 4.03—3.13 (m, C—3H, 22H₂).
- 25 Example 13 9,10-seco-3β-hydroxy-20(S)-hydroxymethyl-pregna-5(E),7(E),10(19)-triene Method A

The phthalazine adduct from Example 3(1) (200 mg) was treated with hydrazine, followed by oxidation as described in the general procedure to give the title product (105 mg; 85%).

## Method B

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The phthalazine adduct from Example 3(3) (250 mg) was similarly converted to give the t-butyldimethylsilyl ether of the title product (166 mg, 90%). This material in refluxing THF (10 ml) was stirred with n-Bu<sub>3</sub>NF (1 M soln. in THF, 2 ml) for 1 hr. Dilution with CH<sub>2</sub>Cl<sub>2</sub>, followed by aqueous work-up and purification by plc gave (267c) (107 mg, 87%).

#### Method C

The product of Method B (160 mg) obtained via the corresponding SO<sub>2</sub> adducts was sillarly converted to the title compound.

## Method D

The triethylsilyl ether of the title compound (160 mg) obtained via the corresponding SO<sub>2</sub> adducts in THF (10 ml) was stirred at room temperature with n-Bu<sub>4</sub>NF (1 M soln. in THF, 2 ml). After about 30 min, the reaction was worked up as for (B) above, to give (267c) (101 mg, 85%).

Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/hexane. m.p.  $104-106^{\circ}$ C; [ $\alpha$ ]<sub>D</sub> =  $+190^{\circ}$  (C = 0.37); UV  $\lambda$ max 273 nm (22640); <sup>1</sup>Hnmr  $\delta$  6.5 and 5.83 (ABq, J=11Hz, C—6H, 7H), 4.93 (s, C—19H), 4.62 (s, C—19H), 4.08—3.12 (m, C—3H, 22H<sub>2</sub>), 1.05 (d, J=5Hz, C—21H<sub>3</sub>), 0.58 (s, C—18H<sub>3</sub>); IR vmax 3450 (s), 2980 (s), 2950 (sh), 1635 (w), 1450 (m), 1050 (s), 1030 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 330; (analysis found: % C, 79.46; H, 9.94; C<sub>22</sub>H<sub>34</sub>O<sub>2</sub> requires % C, 79.95; H, 10.37).

#### Example 14

 $6(R),19-[N,N'-phthalhydrazido]-9,10-seco-3\beta-acetoxy-20(S)-[p-toluenesulphonyloxymethyl]-pregna-5(10),7(E)-diene$ 

The alcohol from Example 3(1) (2.275 g) in pyridine was stirred overnight with p-toluenesulphonyl-chloride (6.25 g) at room temperature. Water was added to the ice cooled mixture and after about 20 min, the mixture extracted with  $CH_2CI_2$ . Acid work-up followed by crystallisation from  $CH_2CI_2$ /ether gave 2.5 g (85%) of the required tosylate (216). m.p. 91—92°C; [ $\alpha$ ]<sub>D</sub> = +308° (c = 1.26); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl; and d, J=7Hz, 2H, tosyl), 7.33 (d, J=7Hz, 2H, tosyl), 5.85 (d, J=10Hz, C—7H), 5.06 (m, C—3H), 4.78 and 4.2 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 3.8 (m, W=12Hz, C—22H<sub>2</sub>), 2.43 (s, t syl), 2.03 (s. OAc), 0.88 (d, J=5Hz, C—21H<sub>3</sub>), 0.13 (s, C—18H<sub>3</sub>); IR vmax 2950 (s), 2900 (sh), 1750 (s), 1650 (s), 1610 (s), 1475 (s), 1240 (s), 1175 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 686; (analysis found: % C, 68.09; H, 6.844; N, 4.00; S, 4.90;  $C_{39}H_{48}O_7N_2S$  requires: % C, 68.19; H, 6.75; N, 4.08; S, 4.67).

#### Example 15

3-m thyi-1-butyn-3-yl tetrahydropyranyl eth r

3-Methyl-1-butyl-3-ol (25 ml, 21.7 g), dihydropyran (50 ml) and p-toluen sulphonic acid (5 mg) were mix d tog ther at 0°C for 1 hr, and then stirr d at room temperature for a further 40 hr. The mixture was conc ntrated and the r sidu added to 5% aqueous NaHCO<sub>3</sub> and extracted with benzene. The organic solution was dried to give after distillation 37.3 g (86%) of the title ether. b.p. 47°C/0.8 mm Hg (lit. 30—33°C/0.5 mm<sup>50</sup>; 57°C/3.5 mm<sup>170</sup>); <sup>1</sup>Hnmr  $\delta$  5.6 (m, THP C2'H), 2.45 (s, C—1H), 1.51 (s, CH<sub>3</sub>), 1.48 (s, CH<sub>3</sub>); IR .(thin film) 3350 (s), 2950 (s), 2900 (sh), 1125 (s), 1070 (s), 1030 (s), cm<sup>-1</sup>.

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#### Example 16

1-mercapto-2-methyl-2-hydroxy-propane

Ethyl-2-mercapto acetate (10 ml) was added to dry ether (150 ml). The well-stirred solution was cooled to 0°C, and an ethereal solution of methyl magnesium bromide (3 M soln., 100 ml., 3.3 eq) was added dropwise over 1.5 hr. The mixture was removed from the ice bath and stirred for an additional 30 min. Ammonium chloride (18 g) in water was carefully added, and the mixture neutralised with hydrochloric acid to form 2 clear layers. The layers were separated and the ether layer washed with water/brine and dried. The solvent was removed under reduced pressure and the product distilled to give 4.4 g of the thiol. b.p. 46°C/16 mm Hg (lit. 64°/26 mm<sup>158a</sup>, 61°/22 mm<sup>158b</sup>); ¹Hnmr δ 2.6 (d, J=9Hz, C—1H<sub>2</sub>), 2.5 (s, exchanges with D<sub>2</sub>O, —OH), 1.38 (t, J=9Hz, —SH), 1.28 (s, 6H, —(CH<sub>3</sub>)<sub>2</sub>); mass spec. m/e 59 (100), 73 (24), 91 (14).

#### Example 17

6(R),19-[N,N'-phthalhydrazido]-23-thia-9,10-seco-3β-acetoxy-25-hydroxy-cholesta-5(10),7(E)-diene

To the tosylate from Example 14 (2.51 g) in THF (125 ml) and HMPTA (3 ml) was added 1-mercapto-2-methylpropan-2-ol from Example 16 (3 ml). The mixture was degassed and NaH (50% dispersion in oil, 1.3 g) was added. After 2 hr, water was added and the mixture diluted with benzene/CH<sub>2</sub>Cl<sub>2</sub>. Acid work-up followed by chromatography and crystallisation from CH<sub>2</sub>Cl<sub>2</sub>/ether gave 1.71 g (77%) of the title sulphide. m.p. 187—188°C;  $[\alpha]_D = +348^\circ$  (c = 0.62); <sup>1</sup>Hnmr  $\delta$  8.3 (m, W=12Hz, 2H, aryl), 7.8 (m, W=10Hz, 2H, aryl), 5.9 (d, J=10Hz, C—7H), 5.07 (m, C—3H), 4.78 and 4.18 (an AB system, J=18Hz, C—19H<sub>2</sub>), 4.75 (d, J=10Hz, C—6H), 2.58 (s, C—24H<sub>2</sub>), 2.03 (s, OAc), 1.23 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 0.98 (d, J=6Hz, C—21H<sub>3</sub>), 0.15 (s, C—18H<sub>3</sub>); IR vmax 3600 (m), 2950 (s), 2900 (sh), 1740 (s), 1640 (s), 1610 (m), 1370 (s), 1350 (s), 1240 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 620; (analysis found: % C, 69.47; H, 7.63; N, 4.43; S, 5.21; C<sub>36</sub>H<sub>48</sub>O<sub>5</sub>N<sub>2</sub>S requires: % C, 69.64; H, 7.79; N, 4.51; S, 5.17).

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#### Example 18

23-thia-9,10-seco-3β,25-dihydroxy-cholesta-5(E),7(E),10(19)-triene

Prepared from the adduct of Example 17 as described in the general procedure as an oil. UV  $\lambda$ max 273 nm;  $^{1}$ Hnmr  $\delta$  6.52 and 5.83 (ABq, J=11Hz, C—6H, 7H), 4.95 (s, C—19H), 4.67 (s, C—19H), 3.85 (m, W=14Hz, C—3H), 2.63 (s, C—24H<sub>2</sub>), 1.3 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.1 (d, W=6Hz, C—21H<sub>3</sub>), 0.58 (s, C—18H<sub>3</sub>).

#### Example 19

3-(3',5'-dinitrobenzoate)-ester

Prepared as described for the compound of Example 6(2) in 67% yield from the adduct of Example 17. Crystalline from ether/hexane. m.p.  $108-110^{\circ}$ C; [a]<sub>0</sub> = +188° (c = 0.742); UV  $\lambda$ max 272 nm (25400) and 262 nm (25400); <sup>1</sup>Hnmr  $\delta$  9.12 (m, 3H, aryl), 6.62 and 5.82 (ABq, J=11Hz, C—6H, 7H), 5.33 (m, W=12Hz, C—3H), 5.00 (s, C—19H), 4.78 (s, C—19H), 2.63 (s, C—24H<sub>2</sub>), 1.27 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.08 (d, W=6Hz, C—21H<sub>3</sub>), 0.45 (s, C—18H<sub>3</sub>); IR vmax 3600 (m), 2950 (s), 2900 (sh), 1740 (s), 1640 (m), 1550 (s), 1350 (s), 1280 (s), 1170 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 612; (analysis found: % C, 64.39; H, 7.26; N, 4.43; S, 5.11; C<sub>33</sub>H<sub>44</sub>O<sub>7</sub>N<sub>2</sub>S requires: % C, 64.68; H, 7.24; N, 4.57; S, 5.23).

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## Example 20

23-thia-9,10-seco-1α-hydroxy-3β,25-bis(triethylsilyloxy)-cholesta-5(E),7(E),10(19)-triene

To the diol of Example 18 (400 mg) in CH<sub>2</sub>Cl (15 ml) was added imidazole followed by triethylsilyl-chloride (450 μl). After 7 hrs, water was added and the mixture diluted with CH<sub>2</sub>Cl<sub>2</sub>. Acid work-up gave the crude bis TES derivative which was used without further purification.

Selenium dioxide (106 mg) was stirred in methanol (5 ml) for 45 min. N-methylmorpholine-N-oxide (NMO) (528 mg) was stirred in  $CH_2Cl_2$  (5 ml) in the presence of anhydrous  $MgSO_4$  for 30 min. The NMO solution was filt r d into a s lution of the crude bis TES derivative in 1,2-dichloro ethane (5 ml) and the mixture warmed to reflux. To this refluxing mixture was add d the  $SeO_2$ /methanol. After 35 min at reflux, the h ating mantle was removed, the mixture diluted with  $CH_2Cl_2$  and washed immediately with 5% aqu ous  $NaHCO_3$  and dri d. Purification by plc gave 233 mg [35% from the adduct of Example 17 of the title 1 $\alpha$ -hydroxy compound as an oil. UV  $\lambda$ max 274 nm;  $^1$ Hnmr  $\delta$  6.58 and 5.92 (ABq, J=12Hz, C—6H, 7H), 5.08 (s, C—19H), 4.97 (s, C—19H), 4.67—4.03 (m, C—1H, 3H), 2.58 (:s, C—24H $_2$ ), 1.32 (s, C—26H $_3$ , 27H $_3$ ).

## Example 21

23-thia-9,10-seco-1α,3β,25-trihydroxy-cholesta-5(E),7(E),10(19)-triene

To the bis TES derivative from Example 20 (112 mg) in THF (5 ml), was added anhydrous tetrabutylammonium fluorid (220 mg) in benzene (3 ml). After 2.25 hr at reflux, the mixture was diluted with ethylacetate, washed with water (3×)/brine and dried. The titl triol (50 mg, 68%), was isolated by plc. Crystalline from CH<sub>2</sub>Cl<sub>2</sub>/hexane. m.p. 129—131°C; [ $\alpha$ l<sub>p</sub> = +184° (c = 0.2175); UV  $\alpha$ max 273 nm (21860); <sup>1</sup>Hnmr  $\alpha$  6.58 and 5.92 (ABq, J=11Hz, C—6H, 7H), 5.12 (s, C—19H), 5.0 (s, C—19H), 4.65—4.0 (m, C—1H, 3H), 2.67 (s, C—24H<sub>2</sub>), 1.28 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.12 (d, J=7Hz, C—21H<sub>3</sub>), 0.57 (s, C—18H<sub>3</sub>); IR vmax 3550 (s), 2950 (s), 2900 (sh), 1640 (w), 1050 (m), 1030 (m), cm<sup>-1</sup>; mass spec. molecular ion m/e = 434; (analysis fund: % C, 71.57; H, 9.57; S, 7.23; C<sub>28</sub>H<sub>42</sub>O<sub>3</sub>S requires: % C, 71.84; H, 9.74; S, 7.38).

#### Example 22

23-thia-9,10-seco-3β,25-dihydroxy-cholesta-5(Z),7(E),10(19)-triene

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A solution of the 5,6-trans-trans vitamin from Example 20 (64 mg) in benzene (30 ml) containing triethylamine (1 drop) and antracene (15 mg) was thoroughly degassed and photoisomerised as described in Example 6(1). The mixture was irradiated for 20 min and the required title vitamin (49 mg, 77%) isolated by plc as an oil. UV  $\lambda$ max 262 nm;  $^{1}$ Hnmr  $\delta$  6.25 and 6.0 (ABq, J=11Hz, C—6H, 7H), 5.03 (s, C—19H), 4.82 (s, C—19H), 3.93 (m, W=18Hz, C—3H), 2.67 (s, C—24H<sub>2</sub>), 1.27 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.08 (d, W=6Hz, C—21H<sub>3</sub>), 0.55 (s, C—18H<sub>2</sub>).

Example 23

The 3-(3′,5′-dinitrobenzoate ester of the product of Example 22 was prepared using the method of Example 6(2). Crystalline from ether/hexane. m.p. 145—148°C;  $[\alpha]_D = +109^\circ$  (C = 0.571); UV  $\lambda$ max shoulders at 260 nm (24900) and 235 nm (30600); ¹Hnmr ŏ 9.08 (m, 3H, aryl), 6.33 and 6.06 (ABq, J=11Hz, C—6H, 7H), 5.33 (m, C—3H), 5.15 (s, C—19H), 4.93 (s, C—19H), 2.65 (s, C—24H2), 1.27 (s, C—26H3, 27H3), 1.08 (d, J=6Hz, C—21H3), 0.57 (s, C—18H3). IR vmax 3750 (m), 2950 (s), 2900 (sh), 1750 (s), 1640 (s), 1550 (s), 1345 (s), 1280 (s), 1170 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 612; (analysis found: % C, 64.7; H, 7.24; O, 18.25; N, 4.36; S, 5.15; C<sub>33</sub>H<sub>44</sub>O<sub>7</sub>N<sub>2</sub>S requires: % C, 64.68; H, 7.24; O, 18.28; N, 4.57; S, 5.23).

## Example 24

23-thia-9,10-seco-1α-hydroxy-3β,25-bis(triethylsilyloxy)-cholesta-5(Z),7(E),10(19)-triene

The corresponding 5(E) compound of Example 20 (180 mg) in benzene (35 ml) containing phenazine (40 mg) and triethylamine (4 drops) was thoroughly degassed and irradiated as described above for 35 min. 137 mg (75%) of the less polar 5(Z) compound was isolated as an oil by plc. UV  $\lambda$ max 263 nm; <sup>1</sup>Hnmr  $\delta$  6.35 and 6.05 (ABq, J=11Hz, C—6H, 7H), 5.27 (s, C—19H), 4.95 (s, C—19H), 4.6—3.93 (m, C—1H, 3H), 2.57 (s, C—24H<sub>2</sub>), 1.2 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>).

#### Example 25

23-thia-9.10-seco-1a,3B,25-trihydroxy-cholesta-5(Z),7(E),10(19)-triene

To the corresponding bis TES derivative from Example 24 (185 mg) in THF (8 ml) was added tetrabutylammonium fluoride (1 M soln. in THF, 2 ml). After 1.25 hr at reflux, the mixture was diluted with  $CH_2Cl_2$ . Aqueous work-up and purification by plc gave 110 mg (90%) of the title triol. Crystalline from ether/hexane. m.p. 124—126°C; [ $\alpha$ ]<sub>D</sub> = +54° (c = 0.37); UV  $\alpha$  264 nm (17400); <sup>1</sup>Hnmr  $\alpha$  6.35 and 6.05 (ABq, J=11Hz, C—6H, 7H), 5.33 (s, C—19H), 5.0 (s, C—19H), 4.65—4.0 (m, C—1H, 3H), 2.63 (s, C—24H<sub>3</sub>), 1.27 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.1 (d, J=6Hz, C—21H<sub>3</sub>), 0.55 (s, C—18H<sub>3</sub>); IR vmax 3550 (s), 2950 (s), 2900 (sh), 1640 (w), 1050 (m), 1030 (m), cm<sup>-1</sup>; mass spec. molecular ion m/e = 434; (analysis found: % C, 71.63; H; 9.61; S, 7.34;  $C_{26}H_{42}O_3S$  requires: % C, 71.84; HJ, 9.74; S, 7.38).

## Example 26

23-thia-9,10-seco-1a-bis(3',5'-dinitrobenzoyloxy)-25-hydroxy-cholesta-5(Z),7(E),10(19)-triene

To the triol from Example 25 (75 mg) in pyridine (3 ml) and benzene (5 ml) was added 3,5-dinitrobenzoylchloride (85 mg). Water was added and the mixture diluted with ether. Work-up as in Example 6(2) and purification by plc gave 97 mg (68%) of the unstable bis (dinitrobenzoate).  $^1$ Hnmr  $\delta$  9.8 (m, 6H, aryl), 6.62 (d, J=11Hz, C—6H), 6.12—5.42 (m, C—1H, 3H, 7H, 19H), 5.32 (s, C—19H), 2.63 (s, C—24H<sub>2</sub>), 1.27 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.08 (broad singlet, C—21H<sub>3</sub>), 0.22 (s, C—18H<sub>3</sub>).

#### . Example 27

23-thia-9,10-seco-1α,3β,25-trihydroxy-cholesta-5(Z),7(E),10(19)-triene-23,S-oxides

The sulphide from Example 25 (100 mg) in methanol (10 ml), ether and water (2 ml) was stirred at room temperature with sodium metaperiodate (50 mg). After 3 hr, a further addition of oxidant (20 mg) was made. After a total of 5 hr, the mixture was diluted with  $CH_2Cl_2$ . Aqueous work-up followed by plc gave 92 mg (89%) of the title sulphoxide mixture. Solid from aceton , methanol/hexane, ether. m.p. 148—155°C; [ $\alpha l_D = +77^{\circ}$  (c = 0.691); UV  $\lambda$ max 263 nm (18150) <sup>1</sup>Hnmr  $\delta$  6.35 and 6.05 (ABq, J=11Hz, C—6H, 7H), 5.33 (s, C—19H), 5.0 (s, C—19H), 4.62—3.72 (m, C—1H, 3H, —OH, exchanges with  $D_2O$ ), 2.83 and 2.75 (br ad singlets, C—24H<sub>2</sub>), 1.52 and 1.38 (C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.23 (broad singlet, C—21H<sub>3</sub>), 0.6 (s, C—18H<sub>3</sub>); IR vmax

3500 (s), 3300 (s), 2950 (s), 2900 (sh), 1620 (w), 1380 (s), 1220 (s), 1070 (s), 1050 (s), 1030 (s), 1600 (s), cm $^{-1}$ ; mass spec. molecular ion m/e = 450; (analysis found: % C, 69.05; H, 9.44; S, 7.13; C<sub>26</sub>H<sub>42</sub>O<sub>4</sub>S requires: % C, 69.29; H, 9.39; S, 7.12).

#### Example 28

23-oxa-9,10-seco-3β,25-dihydroxy-cholesta-5(E),7(E),10(19)-triene

The silyl ether from Example 51 (160 mg) was stirred with n-Bu<sub>4</sub>NF (1 M soln. in THF, 1 ml) in refluxing THF (5 ml) for 40 min. Dilution with  $CH_2Cl_2$ , followed by aqueous work-up and purification by plc gave the title compound (102 mg, 82%). UV  $\lambda$ max 274 nm; <sup>1</sup>Hnmr  $\delta$  6.47 and 5.85 (ABq, J=11Hz, C—6H, 7H), 4.9 (s, C—19H), 4.63 (s, C—19H), 3.83 (m, W=18Hz, C—3H), 3.58—3.07 (m, C—22H<sub>2</sub>), 3.18 (s, C—24H<sub>2</sub>), 1.2 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.03 (d, J=6Hz, C—21H<sub>3</sub>), 0.58 (s, C—18H<sub>3</sub>).

#### Example 29

The 3-(3'-5'-dinitrobenzoate) ester of the product of Example 28 was prepared as described previously for Ex. 6(2). Crystalline from ether/hexane. m.p. 75—77°C;  $[\alpha]_D = +176^\circ$  (c = 0.58); <sup>1</sup>Hnmr  $\delta$  9.15 (m, 3H, aryl), 6.58 and 5.78 (ABq, J=11Hz, C—6H, 7H), 5.3 (m, W=12Hz, C—3H), 5.03 (s, C—19H), 4.73 (s, C—19H), 3.57—3.07 (m, C—22H<sub>2</sub>), 3.2 (s, C—24H<sub>2</sub>), 1.22 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.02 (d, J=6Hz, C—21H<sub>3</sub>), 0.47 (s, C—18H<sub>3</sub>); IR vmax 3500 (m), 2950 (s), 2900 (sh), 1730 (s), 1640 (m), 1550 (s), 1460 (m), 1340 (s), 1270 (s), 1165 (m), cm<sup>-1</sup>; mass spec. molecular ion m/e = 596; (analysis found: % C, 66.31; H, 7.55; N, 4.56; C<sub>33</sub>H<sub>44</sub>O<sub>8</sub>N<sub>2</sub> requires: % C, 66.42; H, 7.43; N, 4.70).

#### Example 30

23-oxa-9,10-seco-3β-(t-butyldimethylsilyloxy)-25-hydroxy-cholesta-5(Z),7(E),10(19)-triene

The corresponding 5(E) compound from Example 51 (160 mg) in benzene (30 ml) and triethylamine (3 drops) containing phenazine (35 mg) was thoroughly degassed and irradiated as described above for 30 min. Purification by plc gave (273a) (138 mg, 86%). UV max = 263 nm;  $^{1}$ Hnmr  $\delta$  6.25 and 6.0 (ABq, J=11Hz, C—6H, 7H), 5.05 (s, C—19H), 4.82 (s, C—19H), 3.92 (m, W=18Hz, C—3H), 3.62—3.10 (m, C—22H<sub>2</sub>), 3.20 (s, C—24H<sub>2</sub>), 1.23 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.03 (d, J=6Hz, C—21H<sub>3</sub>), 0.91 (s, t-Bu), 0.58 (s, C—18H<sub>3</sub>), 0.05 [s, (Si—CH<sub>3</sub>)<sub>2</sub>].

#### Example 31

23-oxa-9,10-seco-3β,25-dihydroxy-cholesta-5(Z),7(E),10(19)-triene

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The corresponding 3-t-butyldimethylsilyl ether from Example 30 (138 mg) was stirred with n-Bu<sub>4</sub>NF (1 M soln. in THF, 2 ml) in refluxing THF (5 ml). After 45 min, the mixture was diluted with CH<sub>2</sub>Cl<sub>2</sub>. Aqueous work-up followed by purification by plc gave the diol (273b) (91 mg, 85%) as an oil. UV  $\lambda$ max 263 nm; <sup>1</sup>Hnmr  $\delta$  6.24 and 6.04 (ABq, J=11Hz, C—6H, 7H), 5.03 (s, C—19H), 4.83 (s, C—19H), 3.92 (m, W=18Hz, C—3H), 3.57—3.12 (m, C—22H<sub>2</sub>), 3.25 (s, C—24H<sub>2</sub>), 1.22 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.03 (d, J=6Hz, C—21H<sub>3</sub>), 0.57 (s, C—18H<sub>3</sub>).

#### Example 32

The 3-(3',5'-dinitrobenzoate) ester of the product of Example 31 was prepared as in Example 6(2). Crystalline from ether/hexane m.p. 136—138°C;  $[\alpha]_D = +101^\circ$  (c = 0.615); <sup>1</sup>Hnmr  $\delta$  9.12 (m, 3H, aryl), 6.22 and 6.01 (ABq, J = 11Hz, C—6H, 7H), 5.23 (m, W=18Hz, C—3H), 5.1 (s, C—19H), 4.92 (s, C—19H), 3.57—3.1 (m, C—22H<sub>2</sub>), 3.2 (s, C—24H<sub>2</sub>), 1.22 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.05 (d, J=6Hz, C—21H<sub>3</sub>), 0.53 (s, C—18H<sub>3</sub>); IR vmax 3550 (m), 2950 (s), 2900 (sh), 1750 (s), 1650 (m), 1555.(s), 1470 (m), 1350 (s), 1280 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 596; (analysis found: % C, 66.34; H, 7.37; N, 4.61; C<sub>33</sub>H<sub>44</sub>O<sub>8</sub>N<sub>2</sub> requires: % C, 66.42; H, 7.43; N, 4.70).

#### Example 33

23-oxa-9,10-seco-3β-(t-butyldimethylsilyloxy)-25-(triethylsilyloxy)-cholesta-5(E),7(E),10(19)-triene

The 25-hydroxy compound from Example 51 (300 mg) in  $CH_2CI_2$  (10 ml) was treated with triethylsilylchloride (130 µl) in the presence of imidazole (200 mg) for 16 hrs. Acid work-up gave the title bis silylated calciferol (274) which was used in the next step without further purification.

#### Example 34

23-oxa-9,10-seco-3α-hydroxy-3β-(t-butyldimethylsilyloxy)-25-(triethylsilyloxy)-cholesta-5(E),7(E),10(19)-triene

Selenium dioxide (60 mg) was stirred in methanol (4 ml) for 45 mins. NMO (300 mg) was stirred in  $CH_2Cl_2$  (4 ml) in the pr sence of anhydrous  $MgSO_4$  for 30 min. The NMO solution was filtered into a solution of the bis silyl ether from Example 33 in 1,2-dichloroethane (4 ml) and the mixture warmed to reflux. To this refluxing mixture was added the  $SeO_2$ /methanol mixture. After 23 min, the h ating mantle was rem ved and the product worked up and isolated as described previously to give 190 mg [51% bas d on Ex. 52] of the titl 1 $\alpha$ -hydroxylat d product. UV  $\alpha$ max 274 nm;  $\alpha$ 1 hnmr  $\alpha$ 3 6.55 and 5.88 (ABq, J=12Hz, C—6H, 7H), 5.1 (s, C—19H), 5.0 (s, C—19H), 4.75—4.02 (m, C—1H, 3H), 3.65—3.12 (m, C—22H<sub>2</sub>), 3.25 (s, C—24H<sub>2</sub>).

### Example 35

23-oxa-9,10-seco-1α,3β,25-trihydroxy-cholesta-5(E),7(E),10(19)-triene

The bis silyl th r fr m Example 34 (190 mg) in THF (6 ml) was refluxed with nBu<sub>4</sub>NF (1 M solution in THF, 2 ml) for 1 hr. The mixture was diluted with CH<sub>2</sub>Cl<sub>2</sub>. Aqueous work-up gave the title triol (103 mg, 84%) aft r purification by plc. Crystalline from chlor form/hexan . mp 141—144°C, [ $\alpha$ ]<sub>D</sub> = +144° (c = 0.355); UV  $\alpha$  x 272 nm (20554); <sup>1</sup>Hnmr (400 MHz)  $\alpha$  6.58 (d, J=12Hz), 5.89 (d, J=12Hz), 5.13 (s, C—19H), 4.98 (s, C—19H), 4.50 (m, W=12Hz, C—1H), 4.26 (m, W=20 Hz, C—3H), 3.43 (m, 1H), 3.30—3.15 (m, C—22H<sub>2</sub>, 24H<sub>2</sub>), 1.20 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.02 (d, J=6Hz, C—21H<sub>3</sub>), 0.58 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 2900 (sh), 1640 (w), 1450 (m), 1380 (m), 1360 (m), 1045 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 418; (analysis found: % C, 74.76; H, 10.33; C<sub>26</sub>H<sub>42</sub>O<sub>4</sub> requires: % C, 74.60; H, 10.11).

#### Example 36

23-oxa-9,10-seco-1α-hydroxy-3β-(t-butyldimethylsilyloxy)-25-(triethylsilyloxy)-cholesta-5(Z),7(E),10(19)-triene

The corresponding 5(E) compound from Example 35 (200 mg) in benzene (35 ml) containing phenazine (40 mg) and triethylamine (4 drops) was irradiated with the hanovia lamp as described previously for 35 min to give, after purification by plc, 155 mg (78%) of the title compound as a less polar, oily product. UV λmax 263 nm; <sup>1</sup>Hnmr δ 6.30 and 6.01 (ABq, J=12Hz, C—6H, 7H), 5.23 (s, C—19H), 4.97 (s, C—19H), 4.67—3.9 (m, C—1H, 3H), 3.53—3.07 (m, C—22H<sub>2</sub>), 3.17 (s, C—24H<sub>2</sub>).

#### Example 37

23-oxa-9,10-seco-1a,3\(\beta\),25-trihydroxy-cholesta-5(Z),7(E),10(19)-triene

The bis silyl ether from Example 36 (155 mg) and n-Bu<sub>4</sub>NF (1 M soln. in THF, 2 ml) were stirred together in refluxing THF (5 ml) for 1 hr. Dilution with  $CH_2Cl_2$  followed by aqueous work-up and purification by plc gave the title triol (252a) (77 mg, 77%). Crystalline from ether/hexane. m.p. 121—123°C; [ $\alpha$ ]<sub>D</sub> = +47° (c = 0.6); UV  $\lambda$ max 264 nm (17200); <sup>1</sup>Hnmr  $\delta$  6.37 and 6.05 (ABq, J=11Hz, C—6H, 7H), 5.33 (s, C—19H), 5.0 (s, C—19H), 4.57—3.87 (m, C—1H, 3H), 3.6—3.1 (m, C—22H<sub>2</sub>), 3.23 (s, C—24H<sub>2</sub>), 1.23 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.05 (d, J=6Hz, C—21H<sub>3</sub>), 0.58 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 2900 (sh), 1640 (w), 1450 (m), 1360 (m), 1045 (s), cm<sup>-1</sup>; mass spec. molecular ion m/e = 418; (analysis found: % C, 74.47; H, 9.97;  $C_{26}H_{42}O_4$  requires: % C, 74.60; H, 10.11).

#### Example 38

9,10-seco-3 $\beta$ -(triethylsilyloxy)-20(S)-(p-toluenesulphonyloxymethyl)-pregna-5(E),7(E),10(19)-triene Method A

To the hydroxy compound from Example 13(D) (400 mg) in pyridine (5 ml) was added tosylchloride (350 mg) and the mixture stirred overnight at room temperature. Water was added and the mixture diluted with ether. Acid work-up gave, after purification by plc, 310 mg (58%) of the title tosylate.

<sup>1</sup>Hnmr δ 7.73 (d, J=8Hz, 2H, aryl), 7.28 (d, J=8Hz, 2H, aryl), 6.43 and 5.81 (ABq, J=11Hz, C—6H, 7H), 4.92 (s, C—19H), 4.63 (s, C—19H), 4.2—3.57 (m, C—3H, 22H<sub>2</sub>), 2.48 (s, aryl-CH<sub>3</sub>); IR vmax (thin film) 2960 (s), 2900 (sh), 1600 (w), 1460 (m), 1360 (s), 1190 (s), 1175 (s), 1090 (s), cm<sup>-1</sup>.

#### Method B

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The crude SO<sub>2</sub> adducts of 9,10-seco-3β-triethylsilyloxy-20(S)-(hydroxymethyl)-pregna-5(E),7(E),10(19)-triene from Example 12(1) (3.2 g) was stirred overnight in pyridine (40 ml) at 5°C with p-toluenesulphonyl chloride (4 g). The mixture was cooled to 0°C, water added and, after a few minutes, the mixture was diluted with Et<sub>2</sub>O. After an acid work-up, the crude oily product (281) was taken up in ethanol (100 ml) and refluxed in the presence of NaHCO<sub>3</sub> (4 g) for 1 hr. The mixture was concentrated and partitioned between CH<sub>2</sub>Cl<sub>2</sub>/ water/brine. The organic solution was dried and chromatographed to give 2.64 g (70%) of the required vitamin (278c) nmr and IR identical to the product obtained by Method A.

## Example 39

9,10-seco-3β-hydroxy-20(S)-[fluoromethyl]-pregna-5(E),7(E),10(19)-triene

The tosylate from Example 38 (200 mg) in THF (5 ml) was refluxed for 45 min in the presence of n-Bu<sub>4</sub>NF (1 M soln. in THF, 1 ml). The mixture was diluted with  $CH_2Cl_2$ . Aqueous work-up followed by purification by plc gave 70 mg (63%) of the title fluoride (279). <sup>1</sup>Hnmr  $\delta$  6.5 and 5.83 (ABq, J=11Hz, C—6H, 7H), 4.97 (s, C—19H), 4.7 (br, s, C—19H, 22H), 4.2—3.6 (m, C—3H, 22H), 1.1 (d, J=6Hz, C 21H<sub>3</sub>), 0.6 (s, C—18H<sub>2</sub>).

#### Example 40

9,10-seco-1α-hydroxy-3β-(triethylsilyloxy)-20(S)-(p-toluenesulphonyloxymethyl)-pregna-5(E),7(E),10(19)-triene

S lenium dioxid (56 mg) was stirr d in acetonitrile (3.5 ml) for 45 min. NMO (280 mg) was stirred in  $CH_2CI_2$  (3.5 ml) in the presence of anhydrous MgSO<sub>4</sub> for 30 min. The NMO solution was filtered into a solution of the 1-desoxy compound from Example 387 (308 mg) in 1,2-dichloroethane (3.5 ml) and the mixture warmed to reflux. To this was added the  $SeO_2/CH_2CN$  mixture, and refluxing continued for a further

5.5 min. The reaction mixture was cooled in an ice bath, diluted with  $CH_2CI_2$  and worked up as previously to give 180 mg (57%) of the title 1-hydroxy compound. <sup>1</sup>Hnmr  $\delta$  7.73 (d, J=8Hz, 2H, aryl), 7.28 (d, J=8Hz, 2H, aryl), 6.43 and 5.81 (ABq, J=11Hz, C—6H, 7H), 5.03 (s, C—19H), 4.93 (s, C—19H), 4.63—3.6 (m, C——1H, 3H, 22H<sub>2</sub>), 2.48 (s, aryl-CH<sub>3</sub>).

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#### Example 41

9,10-seco-1a,3β-dihydroxy-20(S)-(p-toluenesulphonyloxymethyl)-pregna-5(E),7(E),10(19)-triene

The 3-triethylsilylether derivative from Example 40 (180 mg) in THF (5 ml) containing n-Bu<sub>4</sub>NF (1 M soln. in THF, 0.4 ml) was stirred for 15 min. The mixture was diluted with  $CH_2Cl_2$ . An aqueous work-up and purification by plc gave 118 mg (81%) of the title diol. Solid from  $CH_2Cl_2$ /hexane. m.p. 97—99°C; [ $\alpha$ ]<sub>0</sub> = +132° (c = 0.57); UV  $\lambda$ max 272 nm (23360) and 218 nm (15920); <sup>1</sup>Hnmr  $\delta$  7.73 (d, J=8Hz, 2H, aryl), 7.28 (d, J=8Hz, 2H, aryl), 6.43 and 5.81 (ABq, J=11Hz, C—6H, 7H), 5.03 (s, C—19H), 4.93 (s, C—19H), 4.63—3.53 (m, C—1H, 3H, 22H<sub>2</sub>), 2.5 (s, aryl-CH<sub>3</sub>), 1.02 (d, J=6Hz, C—21H<sub>3</sub>), 0.57 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 1600 (w), 1450 (m), 1355 (s), 1190 (s), 1175 (s), cm<sup>-1</sup>.

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#### Example 42

9,10-seco-1a-hydroxy-3β-(triethylsilyoxy)-20(S)-(p-toluenesulphonyloxymethyl)-pregna-5(Z),7(E),10(19)-triene

The corresponding 5(E) compound from Example 40 (225 mg) in benzene (35 ml) containing triethyl20 amine (3 drops) was irradiated as described above with anthracene (45 mg) as triplet sensitizer for 30 min
to give, after plc, 185 mg (82%) of the title compound. UV λmax 263 nm and 216 nm; <sup>1</sup>Hnmr δ 7.73 (d,
J=8Hz, 2H, aryl), 7.3 (d, J=8Hz, 2H, aryl), 6.28 and 5.98 (ABq, J=11Hz, C—6H, 7H), 5.28 (s, C—19H), 4.92 (s,
C—19H), 4.55—3.58 (m, C—1H, 3H, 22H<sub>2</sub>), 2.45 (s, aryl-CH<sub>3</sub>).

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## Example 43

9,10-seco-1α,3β-dihydroxy-20(S)-(p-toluenesulphonyloxymethyl)-pregna-5(Z),7(E),10(19)-triene

The silyl ether from Example 43 (185 mg) in THF (5 ml) containing n-Bu<sub>4</sub>NF (1 M soln. in THF, 0.32 ml) was stirred for 15 min at room temperature. Dilution with  $CH_2CI_2$  aqueous work-up and purification by plc gave the title diol (110 mg, 73%). UV  $\lambda$ max 263 nm (17427) and 216 nm (18672); <sup>1</sup>Hnmr  $\delta$  7.68 (d, J=8Hz, 2H, aryl), 7.23 (d, J=8Hz, 2H, aryl), 6.28 and 5.97 (ABq, J=11Hz, C—6H, 7H), 5.27 (s, C—19H), 4.93 (s, C—19H), 4.57—3.6 (m, C—1H, 3H, 22H<sub>2</sub>), 2.45 (s, aryl-CH<sub>3</sub>), 1.05 (d, J=6Hz, C—21H<sub>3</sub>), 0.52 (s, C—18H<sub>3</sub>).

## Example 44

1-amino-2-methyl-2-hydroxy-propane

To a well-stirred mixture of lithium aluminium hydride (12 g) in ether (200 ml) at 0°C was added dropwise over 1 hr a solution of acetone cyanohydrin (11.2 g, 12 ml) in ether (50 ml). The mixture was stirred at room temperature overnight. After cooling to 0°C, water (24 ml) was cautiously added dropwise. After the quenching was complete, anhydrous Na<sub>2</sub>SO<sub>4</sub> (65 g) was added and stirring at room temperature was continued for a further 2.5 hr. The solid was filtered off and the ether evaporated to give, after distillation, 4.8 g (41%) of the title compound as a viscous, colourless liquid.

b.p. 74—76°C/14 mm Hg (lit.<sup>169</sup> 62—64°C/13 mm Hg)  $n_0^{60} = 1.4463$  (lit.<sup>169</sup>  $n_0^{20} = 1.4467$ ); <sup>1</sup>Hnmr  $\delta$  2.6 (s, 2.6 (s, 2H), 1.87 (s, 3H, exchanges with D<sub>2</sub>O), 1.2 (s, 6H); IR vmax (thin film) 3400 (s), 3000 (m), 1600 (m), 1475 (m), 1380 (m), 1360 (m), 1220 (m), 1170 (m), 1110 (m), 960 (m), cm<sup>-1</sup>.

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## Example 45

23-aza-9,10-seco-1α,3β,25-trihydroxy-cholesta-5(Z),7(E),10(19)-triene

A solution of the toxylate from Example 44 (100 mg) in 1-amino-2-methyl-2-hydroxy-propane (0.5 ml) was degassed and then stirred under argon at 50—55°C for 6 hr and then at room temperature for a further 12 hr. The solution was diluted with CH<sub>2</sub>Cl<sub>2</sub> and washed with water/brine and dried to give, after purification by plc, 44 mg (53%) of the title triol. [ $\alpha$ ]<sub>D</sub> = +24°; UV  $\alpha$  264 nm (15400); <sup>1</sup>Hnmr  $\alpha$  6.38 and 6.07 (ABq, J=11Hz, C—6H, 7H), 5.35 (s, C—19H), 5.02 (s, C—19H), 4.67—3.93 (m, C—1H, 3H), 2.5 (s, C—24H<sub>2</sub>), 1.2 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.02 (d, J=6Hz, C—21H<sub>3</sub>), 0.57 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 2900 (sh), 1640 (w), 1460 (m), 1380 (m), 1055 (m), cm<sup>-1</sup>; mass measurement found: 417.3242; C<sub>26</sub>H<sub>49</sub>O<sub>3</sub>N requires: 417.3243.

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## Example 46

23-aza-9,10-seco-1α,3β,25-trihydroxy-cholesta-5(Z),7(E),10(19)-triene-23-N-acetyl

The crude amine from Example 45 derived from the tosylate (100 mg) as described above, in methanol (5 ml) containing  $K_2CO_3$  (500 mg) was treated with acetic anhydride (0.2 ml). The mixture was diluted with  $CH_2CI_2$  washed with brine and dried to give, aft r plc, 50 mg [55% fr m tosylate] of the title amide. Solid from  $CH_2CI_2$ /hexane. m.p. 107—109°C; [ $\alpha$ ]<sub>D</sub> = -14° (c = 0.49); UV  $\lambda$ max 263 nm (16275);  $^1$ Hnmr  $\delta$  6.37 and 6.05 (ABq, J=11Hz, C—6H, 7H), 5.33 (s, C—19H), 5.0 (s, C—19H), 4.65—4.02 (m, C—1H, 3H), 3.4 (s, C—24H<sub>2</sub>), 2.17 (s, acetyl), 1.22 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 0.95 (d, J=7Hz, C—21H<sub>3</sub>), 0.6 (s, C—18H<sub>3</sub>); IR vmax 3550 (s), 2950 (s), 2900 (sh), 1640 (s), 1460 (m), 1380 (m), 1055 (m), cm<sup>-1</sup>; (analysis found: % C, 70.80; H, 10.12; N, 2.77;  $C_{28}H_{45}O_4N$  requires: C, 73.16; H, 9.87; N, 3.05;  $C_{28}H_{45}O_4N$  .  $H_2O$  requires: % C, 70.40; H, 9.92; N, 2.93).

## - Example 47

9,10-s co-3β,25-dihydroxy-cholesta-5(E),7(E),10(19)-triene

Magnesium turning were washed with diluted HCl/water/acetone/ether and dried in vacuo for 24 hr. The 1-bromo-4-methyl-4-triethylsilylbutane (1 g) in freshly distilled (from LiAlH<sub>4</sub>) THF (10 ml) containing magnesium metal (82 mg) was refluxed for 2 hr.

Cuprous iodide (100 mg) was placed in a flask and purged with argon, whilst cooling to 0°C. To this was added the above Grignard solution (5 ml), and the purple coloured mixture stirred for an additional 30 min at 0°C. A solution of the tosylate (278c) (200 mg) in ether (2 ml) was added and the mixture stirred for 40 min at room temperature. Water was added and the reaction mixture extracted with ether. After an acid work-up, the non-polar product was isolated by plc contaminated with large quantities of low molecular weight alkyl residues. This mixture was stirred with n-Bu<sub>4</sub>NF (1 M soln. in THF, 2 ml) in refluxing THF (5 ml) for 2 hr. Dilution with CH<sub>2</sub>Cl<sub>2</sub> followed by aqueous work-up and purification by plc gave 110 mg [82% from tosylate (278c)] of this previously described title diol. The physical and spectral properties of this material were identical in all respects to the product obtained from the phthalazine adduct.

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## Example 48

9,10-seco-3β,25-dihydroxy-cholest-5(Z),7(E),10(19)-triene

The product from Example 47 (100 mg) in benzene (30 ml) and triethylamine (3 drops) containing anthracene (25 mg) was thoroughly degassed and irradiated for 1 hr as described above to give, after purification by plc, the title 5(Z) compound (90 mg, 82%). The physical and spectral properties of this material were identical in all respects to the product obtained via the phthalazine adduct. A mixed melting point determination of this material and an authentic sample, kindly supplied by Roussel Uclaf, Inc. (Romainville, France) was undepressed.

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#### Example 49

9,10-seco-1α,3β-bis(triethylsilyloxy)-20(S)-(p-toluenesulphonyloxymethyl)-pregna-5(Z),7(E),10(19)-triene
The tosylate (276b) (105 mg) in CH<sub>2</sub>Cl<sub>2</sub> (5 ml) containing imidazole (75 mg) and triethylsilylchloride
(45 μl) was stirred at room temperature for about 15 min. Water was added and the mixture diluted with
CH<sub>2</sub>Cl<sub>2</sub>. Acid work-up gave the non-polar title bis silyl ether which was used without further purification.

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#### Example 50

9,10-seco-1a,3\(\beta\),25-trihydroxy-cholesta-5(Z),7(E),10(19)-triene

To the alkyl copper reagent at 0°C prepared exactly as described above, was added a solution of the above tosylate (276c) in THF (3 ml) and the mixture stirred at room temperature for 25 min. Work-up and purification as in Example 6(1) gave the tris triethylsilyl derivative contaminated with large quantities of low molecular weight alkyl residues. This mixture was treated with n-Bu<sub>4</sub>NF (1 M soln. in THF, 4 ml) in THF (5 ml) for 20 min at room temperature followed by 1.5 hr at reflux to give, after the usual work-up and purification by plc, a mixture of the title steroidal triol [(38 mg, 63% from (276b)] contaminated with isopentane diol (10 mg). Dissolution of this mixture in CHCl<sub>3</sub> gave the required product as its crystalline CHCl<sub>3</sub> solvate (25 mg). m.p. 99—105°C (lit. 106—112°C<sup>142</sup>, 103—106°C<sup>138</sup>); [ $\alpha$ ]<sub>0</sub> (Et<sub>2</sub>O) = +35° (c = 0.86); UV  $\lambda$ max 264 nm (16820); <sup>1</sup>Hnmr  $\delta$  (acetone-d<sub>6</sub>) 8.07 (s, CHCl<sub>3</sub>), 6.35 and 6.18 (ABq, J=12Hz, C—6H, 7H), 5.38 (s, C—19H), 4.93 (:s, C—19H), 4.7—4.07 (m, C—1H, 3H), 1.2 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.0 (broad singlet, C—21H<sub>3</sub>), 0.6 (s, C—18H<sub>3</sub>); IR vmax 3500 (s), 2950 (s), 2900 (sh), 1640 (w), 1480 (m), 1440 (m), 1380 (m), 1360 (m), 1140 (m), 1050 (s).

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#### Example 51

23-oxa-9,10-seco-3B-(t-butyldimethylsilyloxy)-25-hydroxy-cholesta-5(E),7(E),10(19)-triene

The 22-hydroxy compound (167b) (425 mg) in benzene (5 ml) was refluxed with isobutylene epoxide (1 ml) in the presence of dibenzo-18-crown-6 (100 mg) and potassium t-butoxide (500 mg) for 55 min. Water was added and the mixture diluted with  $CH_2CI_2$ . The organic layer was washed with aqueous  $K_3PO_4$ /water/5% aqueous NaHCO<sub>3</sub>/brine and dried. Purification by plc gave 330 mg (67%) of the slightly less polar oily product. <sup>1</sup>Hnmr  $\delta$  6.45 and 5.85 (ABq, J=12Hz, C—6H, 7H), 4.9 (s, C—19H), 4.63 (s, C—19H), 3.92 (m, W=18Hz, C—3H), 3.63—3.12 (m, C—22H<sub>2</sub>), 3.22 (s, C—24H<sub>2</sub>), 1.23 (s, C—26H<sub>3</sub>, 27H<sub>3</sub>), 1.05 (d, J=6Hz, C—21H<sub>3</sub>), 0.92 (s, t-Bu), 0.6 (s, C—18H<sub>3</sub>), 0.05 s, [(Si—CH<sub>3</sub>)<sub>2</sub>].

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#### Claims for the Contracting States: BE CH DE FR GB IT LI LU NL SE

## 1. Compounds of the general formula

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wherein R represents a hydrogen atom or a hydroxyl protecting group, Y represents a hydrogen atom of an optionally protected hydroxyl group, X represents —SO<sub>2</sub> or the residue of a diacylazo dienophile and either R1 represents a halogen atom a hydrocarbylsulphonyloxy group or a group of the formula -Z-R3 (in which Z represents —O—, —S—, —SO—, —NR<sup>4</sup>— or —CR<sup>4</sup>R<sup>5</sup>— and R<sup>3</sup>, R<sup>4</sup> and R<sup>5</sup>, which may be the same or different, each represent a hydrogen atom or a straight or branched aliphatic group having 1—12 carbon atoms and which may optionally carry one or more substituents) and R2 represents a hydrogen atom or R1 and R2 together represent an oxo group or an optionally substituted alkylidene group, except that R1 and R2 together with the group -CH(CH<sub>3</sub>)CH- to which they are attached do not represent a group having the branched 17 $\beta$ -hydrocarbyl side chain skeleton of vitamin  $D_2$  or vitamin  $D_3$ .

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2. Compounds as claimed in claim 1 in which the dienophile is a cyclic diacylazo compound.

3. Compounds of general formula IV or IVA,

40 45 50 RO 55 IVa (cis) IV (trans)

wherein R, Y, R<sup>1</sup> and R<sup>2</sup> are as defined in claim 1.

4. Compounds of general formulae I, IV or IVa as claimed in any one of claims 1 to 3 wherein R1 50 r pr sents a halogen atom, a hydr xyl or tosyloxy gr up or a group of formula:

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(in which Z' represents —O—, —S—, —NH— or —SO— and R<sup>6</sup> represents a hydrogen at m or a hydroxyl protecting group) and R<sup>2</sup> r pr sents a hydrogen atom or R<sup>1</sup> and R<sup>2</sup> tog ther represent an alkylidene group having 1 to 8 carbon atoms optionally substituted by one or more substituents selected from halog n atoms and optionally protected hydroxyl groups.

5. A pr c ss for the preparation f c mpounds f general formula I as defined in claim 1 in which R<sup>1</sup> and R<sup>2</sup> together repres nt an oxo group which comprises subjecting a compound of formula III.

(wherein R, Y and X are as defined in claim 1) to oxidative cleavage.

6. A process as claimed in claim 5 wherein the aldehyde of formula I so formed is subsequently reduced to give a compound of formula I wherein R<sup>1</sup> represents a hydroxyl group.

7. A process as claimed in claim 5 wherein the aldehyde of formula I so formed is subsequently reacted with a Wittig reagent to give a compound of formula I wherein R<sup>1</sup> and R<sup>2</sup> together represent an optionally substituted alkylidene group, the double bond of which may then, if desired, be reduced.

8. A process for the preparation of compounds of general formula IV or IVa as defined in claim 3 which comprises deprotecting a corresponding compound of formula I as defined in claim 1 by removal of the residue X and subsequently, optionally after conversion of the group R<sup>1</sup> to another group R<sup>1</sup>, isomerising the compound of formula IV thus obtained to a compound of formula IVa.

9. A process as claimed in claim 6 or claim 8 wherein a product is obtained in which R¹ represents a hydroxyl group and the said hydroxyl group is converted into a halogen atom or a hydrocarbyl-sulphonyloxy group.

10. A process as claimed in any one of claims 6, 8 and 9 wherein a product is obtained in which R<sup>1</sup> represents a halogen atom or a hydroxyl or hydrocarbyl sulphonyloxy group which product is converted into a product wherein R<sup>1</sup> represents a group of formula —ZR<sup>3</sup> as defined in claim 1 other than a hydroxyl group.

## Claims for the Contracting State: AT

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1. A process for the preparation of compounds of the general formula

(wherein R represents a hydrogen atom or a hydroxyl protecting group, Y represents a hydrogen atom or an optionally protected hydroxyl group, X r presents—SO<sub>2</sub> or the residue of a diacylazo dienophile and either R<sup>1</sup> represents a halogen atom a hydrocarbylsulphonyloxy group or a group of the formula—Z—R<sup>3</sup>

(in which Z represents -O-, -S-, -SO-,  $-NR^4-$  or  $-CR^4R^5-$  and  $R^3$ ,  $R^4$  and  $R^5$ , which may be the same or different, each r present a hydrog r atom or a straight or branched aliphatic group having r carbon atoms and which may optionally carry one or more substituents) and r represents a hydrogen atom or r and r together represent an oxo group or an optionally substituted alkylidene group, except that r and r together with the group  $-CH(CH_3)CH-$  to which they are attached do not represent a group having the branched r hydrocarbyl side chain skeleton of vitamin r or vitamin r which comprises subjecting a compound of formula III,

(wherein R, Y and X ae as defined above) to oxidative cleavage to form an aldehyde of formula I (in which  $R^1$  and  $R^2$  together represent an oxo group) and subsequently, if desired

either reacting the said aldehyde of formula I with a Wittig reagent to give a compound of formula I wherein R<sup>1</sup> and R<sup>2</sup> together represent an optionally substituted alkylidene group, the double bond of which may then, if desired, be reduced.

or reducing the said aldehyde of formula I to give a compound of formula I wherein R<sup>1</sup> represents a hydroxyl group, the said hydroxyl group being then optionally converted into a halogen atom, a hydrocarbylsulphonyloxy group or a group of formula —ZR<sup>3</sup> as defined above other than a hydroxyl group.

2. A process as claimed in claim 1 in which the dienophile is a cyclic diacylazo compound.

3. A process for the preparation of compounds of general formula IV or IVa,

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(wher in R, Y, R<sup>1</sup> and R<sup>2</sup> are as defin d in claim 1) which c mprises deprotecting a corresponding compound of formula I as defined in claim 1 by rem val of the residue X and subsequently, opti nally aft r conversion of the gr up R<sup>1</sup> to another group R<sup>1</sup>, isomerising the compound of formula IV thus obtained to a compound of formula IVa.

4. A process as claimed in any preceding claim f r the preparation of compounds of general formula I, IV or IVa wherein R<sup>1</sup> represents a halogen atom, a hydroxyl or tosyloxy group or a group of formula:

(in which Z' represents —O—, —S—, —NH— or —SO— and R<sup>6</sup> represents a hydrogen atom or a hydroxyl protecting group) and R<sup>2</sup> represents a hydrogen atom or R<sup>1</sup> and R<sup>2</sup> together represent an alkylidene group having 1 to 8 carbon atoms optionally substituted by one or more substituents selected from halogen atoms and optionally protected hydroxyl groups.

5. A process as claimed in any preceding claim wherein a product is obtained in which R<sup>1</sup> represents a hydroxyl group and the said hydroxyl group is converted into a halogen atom or a hydrocarbyl-sulphonyloxy group.

6. A process as claimed in any preceding claim wherein a product is obtained in which R<sup>1</sup> represents a halogen atom or a hydroxyl or hydrocarbylsulphonyloxy group which product is converted into a product wherein R<sup>1</sup> represents a group of formula —ZR<sup>3</sup> as defined in claim 1 other than a hydroxyl group.

## Patentansprüche für die Vertragsstaaten: BE CH DE FR GB IT LI LU NL SE

## 1. Verbindungen der allgemeinen Formel

worin R ein Wasserstoffatom oder eine Hydroxyl-Schutzgruppe bedeutet, Y ein Wasserstoffatom oder eine gegebenenfalls geschützte Hydroxylgruppe darstellt, X für —SO<sub>2</sub> oder den Rest eines Diacylazo-Dienophils steht und entweder R¹ bedeutet ein Halogenatom oder eine Hydrocarbylsulfonyloxygruppe oder eine Gruppe der Formel —Z—R³ (worin Z für —O—, —S—, —SO—, —NR⁴— oder —CR⁴R⁵— steht und R³, R⁴ und R⁵, welche gleich oder verschieden sein können, jeweils ein Wasserstoffatom oder eine gerade oder verzweigte, aliphatische Gruppe mit 1 bis 12 Kohlenstoffatomen bedeuten und die gegebenenfalls einen oder mehrere Substituenten tragen kann) und R² ein Wasserstoffatom bedeutet oder R¹ und R² bedeuten zusammen eine Oxogruppe oder eine gegebenenfalls substituierte Alkylidengruppe, mit der Ausnahme, daß R¹ und R² zusammen mit der Gruppe —CH(CH₃)CH—, an die sie gebunden sind, nicht eine Gruppe bedeuten, welche das verzweigte 17β-Hydrocarbyl-Seitenkettengerüst von Vitamin D₂ oder Vitamin D₃ hat.

2. Verbindungen gemäß Anspruch 1, worin das Dienophil eine cyclische Diacylazo-Verbindung ist.

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## 3. Verbindungen d r allgemeinen Formel IV oder IVa

worin R, Y, R<sup>1</sup> und R<sup>2</sup> wie in Anspruch 1 definiert sind.

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4. Verbindungen der allgemeinen Formeln I, IV oder IVa, wie in- einem der Ansprüche 1 bis 3 beansprucht, worin R<sup>1</sup> ein Halogenatom, eine Hydroxyl- oder Tosyloxygruppe oder eine Gruppe der Formel

bedeutet (worin Z' für —O—, —S—, —NH— oder —SO— steht und R<sup>6</sup> ein Wasserstoffatom oder eine Hydroxyl-Schutzgruppe bedeutet) und R<sup>2</sup> ein Wasserstoffatom bedeutet oder R<sup>1</sup> und R<sup>2</sup> bedeuten zusammen eine Alkylidengruppe mit 1 bis 8 Kohlenstoffatomen, gegebenenfalls substituiert durch einen oder mehrere Substituenten, ausgewählt aus Halogenatomen und gegebenenfalls geschützten Hydroxylgruppen.

5. Verfahren zur Herstellung von Verbindungen der allgemeinen Formel I, wie in Anspruch 1 definiert, worin R¹ und R² zusammen eine Oxogruppe bedeuten, dadurch gekennzeichnet, daß eine Verbindung der Formel III

(worin R, Y und X wi in Anspruch 1 definiert sind) der oxidativen Spaltung unterworfen wird.

6. Verfahren gemäß Anspruch 5, dadurch gekennzeichnet, daß der so gebildete Aldehyd der Formel I anschließend reduziert wird, um eine Verbindung der Formel I zu ergeben, worin R¹ eine Hydroxylgruppe darstellt.

7. Verfahren gemäß Anspruch 5, dadurch gekennzeichnet, daß-der so gebildete Aldehyd der Formel I anschließend mit ein m Wittig-Reagens umgesetzt wird, um eine Verbindung der Formel I zu rgeben,

worin R<sup>1</sup> und R<sup>2</sup> zusammen eine gegebenenfalls substituierte Alkylidengruppe darstellen, deren Doppelbindung dann gewünscht nfalls reduziert werd n kann.

8. Verfahren zur Herstellung von Verbindungen der allgemeinen F rmel IV oder IVa, wi in Anspruch 3 definiert, dadurch g kennzeichn t, daß in ein r entsprechend n Verbindung der F rmel I, wi in Anspruch 1 definiert, die Schutzgruppe durch Entfernung des Restes X entfernt wird und anschließend gegebenenfalls nach Überführung der Gruppe R¹ in eine andere Gruppe R¹ die so erhaltene Verbindung der Formel IV zu einer Verbindung der Formel IVa isomerisiert wird.

9. Verfahren gemäß Anspruch 6 oder Anspruch 8, dadurch gekennzeichnet, daß ein Produkt erhalten wird, worin R¹ eine Hydroxylgruppe bedeutet, und diese Hydroxylgruppe in ein Halogenatom oder eine Hydrocarbylsulfonyloxygruppe umgewandelt wird.

10. Verfahren gemäß einem der Ansprüche 6, 8 und 9, dadurch gekennzeichnet, daß ein Produkt erhalten wird, worin R¹ ein Halogenatom oder eine Hydroxyl- oder Hydrocarbylsulfonyloxygruppe darstellt, welches Produkt in ein Produkt überführt wird, worin R¹ eine Gruppe der Formel —ZR³, wie in Anspruch 1 definiert, anders als eine Hydroxylgruppe, darstellt.

## Patentansprüche für den Vertragsstaat: AT

## 1. Verfahren zur Herstellung einer Verbindung der allgemeinen Formel

[worin R ein Wasserstoffatom oder eine Hydroxyl-Schutzgruppe bedeutet, Y ein Wasserstoffatom oder eine gegebenenfalls geschützte Hydroxylgruppe darstellt, X für —SO $_2$  oder den Rest eines Diacylazo-Dienophils steht und entweder R¹ ein Halogenatom oder eine Hydrocarbylsulfonyloxygruppe oder eine Gruppe der Formel —Z—R³ bedeutet (worin Z für —O—, —S—, —SO—, —NR⁴ oder —CR⁴R⁵ steht und R³, R⁴ und R⁵, die gleich oder verschieden sein könne, jeweils ein Wasserstoffatom oder eine gerade oder verzweigte, aliphatische Gruppe mit 1 bis 12 Kohlenstoffatomen bedeuten und die gegebenenfalls einen oder mehrer Substituenten tragen können) und R² ein Wasserstoffatom darstellt oder R¹ und R² bedeuten zusammen eine Oxogruppe oder eine gegebenenfalls substituierte Alkylidengruppe, mit der Ausnahme, daß R¹ und R² zusammen mit der Gruppe —CH(CH $_3$ )CH—, an die sie gebunden sind, nicht eine Gruppe bedeuten, welche das verzweigte 17β-Hydrocarbyl-Seitenkettengerüst von Vitamin D $_2$  oder Vitamin D $_3$  hat], dadurch gekennzeichnet, daß eine Verbindung der Formel III

(worin R, Y und X wie vorstehend definiert sind) der oxidativen Spaltung zur Bildung eines Aldehyds der Formel I (worin R¹ und R² zusammen eine Oxogruppe darstellen) unterworfen wird und anschliß nd gewünschtenfalls

entweder dieser Aldehyd der Formel I mit einem Wittig-Reagens umgesetzt wird, um eine Verbindung der Formel I zu ergeben, worin R¹ und R² zusammen eine geg benenfalls substituierte Alkylidengruppe darstellen, deren Doppelbindung dann gewünschtenfalls reduziert werden kann;

oder der genannte Aldehyd der Formel I reduziert wird, um eine Verbindung der Formel I zu ergeben, worin R¹ eine Hydroxylgruppe darstellt und diese Hydroxylgruppe dann gegebenenfalls in ein Halogenatom, eine Hydrocarbylsulfonyloxygruppe oder eine Gruppe der Formel —ZR³, wie vorstehend definiert, anders als eine Hydroxylgruppe, überführt wird.

2. Verfahren gemäß Anspruch 1, dadurch gekennzeichnet, daß das Dienophil eine cyclische Diacylazo-Verbindung ist.

3. Verfahren zur Herstellung von Verbindungen der allgemeinen Formel IV oder IVa

(worin R, Y, R<sup>1</sup> und R<sup>2</sup> wie in Anspruch 1 definiert sind), dadurch gekennzeichnet, daß in einer entsprechenden Verbindung der Formel I, wie in Anspruch 1 definiert, die Schutzgruppe durch Entfernung des Restes X entfernt wird und anschließend, gegebenenfalls nach Umwandlung der Gruppe R<sup>1</sup> in eine andere Gruppe R<sup>1</sup>, die so erhaltene Verbindung der Formel IV zu einer Verbindung der Formel IVa isomerisiert wird.

4. Verfahren gemäß einem der vorhergehenden Ansprüche zur Herstellung von Verbindungen der allgemeinen Formeln I, IV oder IVa, worin R¹ eine Halogenatom, eine Hydroxyl- oder Tosyloxygruppe oder eine Gruppe der Formel

bedeutet (worin Z' für —O—, —S—, —NH— oder —SO— steht und R<sup>6</sup> ein Wasserstoffatom oder eine Hydroxyl-Schutzgruppe darstellt) und R<sup>2</sup> ein Wasserstoffatom bedeutet oder R<sup>1</sup> und R<sup>2</sup> zusammen eine Alkylidengruppe mit 1 bis 8 Kohlenstoffatomen, gegebenenfalls durch einen oder mehrere Substituenten, ausgewählt aus Halogenatomen und gegebenenfalls geschützten Hydroxylgruppen, substituiert, darstellen.

5. Verfahren gemäß einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß ein Produkt erhalten wird, worin R¹ eine Hydroxylgruppe darstellt und diese Hydroxylgruppe in ein Halogenatom oder eine Hydrocarbylsulfonyloxygruppe überführt wird.

6. Verfahren gemäß einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß ein Produkt erhalten wird, worin R¹ ein Halogenatom oder eine Hydroxyl- oder Hydrocarbylsulfonyloxygruppe bedeutet, w Iches Produkt in ein Produkt überführt wird, worin R¹ eine Gruppe der Form I —ZR³, wie in Anspruch 1 definiert, anders als ein Hydr xylgruppe, darstellt.

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# R vendications pour les Etats contractants: BE CH DE FR GB IT LI LU NL SE

#### 1. Compos's répondant à la formul g'nérale

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H<sub>2</sub>X

dans laquelle R représente un atome d'hydrogène ou un groupe protégeant la fonction hydroxyle, Y représente un atome d'hydrogène ou un radical hydroxyle éventuellement protégé, X représente un radical —SO<sub>2</sub> ou le reste d'un diacylazodiénophile et R¹ représente un atome d'halogène, un radical hydrocarbylsulfonyloxy ou un groupe de la formule —Z—R³ (où Z représente —O—, —S—, —SO—, —NR⁴— ou —CR⁴R⁵— et R³, R⁴ et R⁵, qui peuvent être identiques ou différents, représentent chacun un atom d'hydrogène ou un radical aliphatique à chaîne droite ou à chaîne ramifiée, possédant de 1 à 12 atomes de carbone et qui peut éventuellement porter un ou plusieurs substituants) et R² représente un atome d'hydrogène, ou bien R¹ et R² représentent ensemble un radical oxo ou un groupe alkylidèn éventuellement substitué, à l'exception que R¹ et R² ne forment pas, ensemble avec le groupe —CH(CH₃)CH— auquel ils sont attachés, un radical possédant le squelette de la chaîne latérale 17β-hydrocarbylique ramifiée de la vitamine D₂ ou de la vitamine D₃.

2. Composés suivant la revendication 1, caractérisés en ce que le diénophile est un composé diacylazoïque cyclique.

3. Composés des formules générales IV et IVa

dans I squelles R, Y, R<sup>1</sup> et R<sup>2</sup> possèdent les significati ns qui leur ont été précédemment attribuées dans la revendication 1.

4. Composés des formules générales I, IV et IVa, suivant l'un quelconque des revendications 1 à 3, dans lesquilles R¹ repr´sente un atome d'halog`n , un radical hydrixyle ou tosyloxy, ou un groupe de la firmule

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(dans laquelle Z' représente —O—, —S—, —NH— ou —SO— et R<sup>6</sup> représente un atome d'hydrogène ou un radical protégeant la fonction hydroxyle) et R<sup>2</sup> représente un atome d'hydrogène, ou bien R<sup>1</sup> et R<sup>2</sup> représentent ensemble un groupe alkylidène possédant de 1 à 8 atomes de carbone, éventuellement substitué par un ou plusieurs substituants choisis parmi les atomes d'halogènes et les radicaux hydroxyle éventuellement protégés.

5. Procédé de préparation de composés de la formule générale I suivant la revendication 1, dans laquelle R¹ et R² représentent ensemble un groupe oxo, caractérisé en ce que l'on soumet un composé de la formula III

(dans laquelle R, Y et X possèdent les significations qui leur ont été précédemment attributées dans la revendication 1) à une scission oxydante.

6. Procédé suivant la revendication 5, caractérisé en ce que l'on réduit ensuite l'aldéhyde de la formule l ainsi formé de façon à obtenir un composé de la formule l dans laquelle R¹ représente le radical hydroxyle.

7. Procédé suivant la revendication 5, caractérisé en ce que l'on fait ensuite réagir l'aldéhyde de la formule I ainsi formé sur un réactif de Wittig de manière à obtenir un composé de la formule I dans laquelle R¹ et R² représentent ensemble un groupe alkyldène éventuellement substitué dont la double liaison peut ensuite être réduite si on le souhaite.

8. Procédé de préparation de composés de la formule IV on de la formule IVa telles que définies dans la revendication 3, caractérisé en ce que l'on déprotège un composé correspondant de la formule I telle que définie dans la revendication 1 par l'enlèvement du résidu X et ensuite, éventuellement après conversion du groupe R<sup>1</sup> en un autre groupe R<sup>1</sup>, on isomérise le composé de la formule IV ainsi obtenu en un composé de la formule IVa.

9. Procédé suivant la revendication 6 ou la revendication 8, caractérisé en ce que l'on obtient un produit dans lequel R¹ représente un radical hydroxyle et on convertit le radical hydroxyle en question en un atome d'halogène ou un radical hydrocarbylsulfonyloxy.

10. Procédé suivant l'une quelconque des revendications 6, 8 et 9, caractérisé en ce que l'on obtient un produit dans lequel R¹ représente un atome d'halogène ou un radical hydroxyle ou hydrocarbyl-sulfonyloxy, produit que l'on convertit ensuite en une substance dans laquelle R¹ représente un groupe de la formule —ZR³ telle que définie dans la revendication 1, autre qu'un radical hydroxyle.

## Revendications pour l'Etat contractant AT:

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# 1. Proc'dé de préparation d composés de la formul générale

(dans laquelle R représente un atome d'hydrogène ou un groupe protégeant la fonction hydroxyle, Y 25 - représente un atome d'hydrogène ou un radical hydroxyle éventuellement protégé, X représente un radical —SO₂ ou le reste d'un diacylazodiénophile et R¹ représente un atome d'halogène, un radical hydrocarbylsulfonyloxy ou un groupe de la formule —Z—R³ (où Z représente —O—, —SO—, —SO—, -NR<sup>4</sup>— ou —CR<sup>4</sup>R<sup>5</sup>— et R<sup>3</sup>, R<sup>4</sup> et R<sup>5</sup>, qui peuvent être identiques ou différents, représentent chacun un atome d'hydrogène ou un radical aliphatique à chaîne droite ou à chaîne ramifiée, possédant de 1 à 12 atomes de carbone et qui peut éventuellement porter un ou plusieurs substituants) et R2 représente un atome d'hydrogène, ou bien R1 et R2 représentent ensemble un radical oxo ou un groupe alkylidène éventuellement substitué, à l'exception que R1 et R2 ne forment pas, ensemble avec le groupe --CH(CH<sub>3</sub>)CH-- auquel ils sont attachés, un radical possédant le squelette de la chaîne latérale 17βhydrocarbylique ramifiée de la vitamine D2 ou de la vitamine D3) caractérisé en ce que l'on soumet un composé de la formule III

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(dans laquelle R, Y et X possèdent les significations qui leur ont été précédemment attribuées) à une scission oxydante de manière à former un aldéhyde de la formule I (dans laquelle R1 et R2 représentent ensemble un radical oxo) et ensuite, si on le souhaite,

on fait réagir l'aldéhyde en question de la formule I sur un réactif de Wittig de manière à obtenir un composé de la formule I dans laquelle R1 et R2 représentent ensemble un radical alkylidène éventuellement substitué dont on peut ensuite réduire la double liaison si on le souhaite, ou bien on réduit l'aldéhyde en question de la formule I de façon à obtenir un composé de la formule I dans laquelle R1 représent un radical hydroxyl , le radical hydroxyle étant ensuite éventuellement converti en un atome d'halogène, un radical hydrocarbylsulfonyloxy ou un groupe de la formule -- ZR3 t il que pr cédemment définie autre qu'un radical hydr xyle.

2. Procédé suivant la revendication 1, caractérisé en ce que le diénophile est un composé diacylazoïque cyclique.

3. Procédé de préparati n de composés de la formule général IV ou de la f rmule générale IVa

(dans lesquelles R, Y, R<sup>1</sup> et R<sup>2</sup> possédent les significations qui leur ont été précédemment attributées dans la revendication 1), caractérisé en ce que l'on déprotège un composé correspondant de la formule I telle que définie dans la revendication 1 par l'enlèvement du reste X et ensuite, éventuellement après

conversion du groupe R<sup>1</sup> en un autre groupe R<sup>1</sup>, on isomérise le composé de la formule IV ainsi obtenu en un composé de la formule IVa.

4. Procédé suivant l'une quelconque des revendications précédentes de préparation de composés des formules générales I, IV ou IVa dans lesquelles R¹ représente un atome d'halogène, un radical hydroxyle ou tosyloxy, ou un groupe de la formule

40 (dans laquelle Z' représente —O—, —S—, —NH— ou —SO— et R<sup>6</sup> représente un atome d'hydogène ou un radical protégeant la fonction hydroxyle) et R<sup>2</sup> représente un atome d'hydrogène, ou bien R<sup>1</sup> et R<sup>2</sup> représentent ensemble un radical alkylidène possédant de 1 à 8 atomes de carbone, éventuellement substitué par un ou plusieurs substituants choisis parmi les atomes d'halogène et les radicaux hydroxyle éventuellement protégés.

5. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que l'on obtient un produit dans lequel R¹ représente un radical hydroxyle et on convertit le radical hydroxyle en question en un atome d'halogène ou un radical hydrocarbylsulfonyloxy.

6. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que l'on obtient un produit dans lequel R¹ représente un atome d'halogène ou un radical hydroxyle ou hydrocarbylsulfonyloxy, produit que l'on convertit ensuite en un substance dans laquelle R¹ représente un groupe de la formule —ZR³ telle que définie dans la revendication 1 autre qu'un radical hydroxyle.

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